

Europäisches Pat ntamt
European Patent Office
Offic européen des brev ts



11 Publication number:

0 553 852 A2

(12)

EUROPEAN PATENT APPLICATION

- 21 Application number: 93101413.8
- 2 Date of filing: 29.01.93

(1) Int. Cl.5. **H01L 21/76**, H01L 21/20, H01L 21/322

- Priority: 30.01.92 JP 38458/92
 30.01.92 JP 38460/92
 31.01.92 JP 16513/92
 31.01.92 JP 16523/92
 31.01.92 JP 16524/92
- (3) Date of publication of application: 04.08.93 Bulletin 93/31
- Designated Contracting States:
 DE FR GB

- 71 Applicant: CANON KABUSHIKI KAISHA 30-2, 3-chome, Shimomaruko, Ohta-ku Tokyo(JP)
- Inventor: Sato, Nobuhiko, c/o Canon Kabushiki Kaisha 30-2, 3-chome, Shimomaruko, Ohta-ku Tokyo 146(JP) Inventor: Yonehara, Takao, c/o Canon Kabushiki Kaisha 30-2, 3-chome, Shimomaruko, Ohta-ku Tokyo 146(JP) Inventor: Sakaguchi, Kiyofumi, c/o Canon Kabushiki Kaisha 30-2, 3-chome, Shimomaruko, Ohta-ku Tokyo 146(JP)
- Representative: Tiedtke, Harro, Dipl.-ing. Patentanwälte Tiedtke-Bühling- Kinne & Partner Bavariaring 4 Postfach 20 24 03 W-8000 München 2 (DE)
- Process for producing semiconductor substrate.
- A process for producing a semiconductor substrate comprises heat-treating a substrate having a monocrystal thereon in a reducing atmosphere.

BACKGROUND OF THE INVENTION

Fi Id of the Inv ntion

Th present invention relates to a process for producing a semiconductor substrate, particularly to a process for producing a semiconductor substrat which is applicable to semiconductor devices, integrated circuits, fine mechanical devices, and the like, or which is suitable for dielectric separation, and electronic devices and integrated circuits formed on a monocrystalline semiconductor layer on an insulator.

Related Background Art

In recent years, integrated circuits have come to be remarkably more highly integrated. As the results, higher precision is required in of processing of the substrate surface thereof to meet much sever r conditions. If the surface becomes rough during the processing, the roughness needs to be eliminated for obtaining a flat smooth surface. Contamination with impurity and particles have also to be avoided.

Polishing is generally employed for flattening the surface of a monocrystal. The polishing is conducted such that the surface to be polished is pressed against polishing cloth made of urethane and is rubbed with it while a polishing liquid composed of a neutral solution or an alkaline solution (e.g., KOH) and a polishing particulate material suspended therein is dropped thereto, to abrade and flatten the surface. For example, a monocrystalline silicon surface is polished and flattened by dropping a suspension, in an alkaline solution (e.g., KOH), of colloidal silica composed of silicon oxide as polishing particulate material.

With the modern technique of polishing of semiconductor substrates, practicably achievable lev I of surface roughness is several nm in terms of p-v (maximum-minimum difference) and about 0.1 nm in terms of rms (mean square deviation).

Another technique of flattening is heat-treatm nt.

S. Nakashima and K. Izumi reported that the roughness of the surface having a great number of concavities of several tens of nm was eliminated by heat-tr atment at 1260 °C in a nitrogen atmosphere for 2 hours, or at 1300 °C in an argon atmosphere containing 0.5% oxygen for 4 hours, but the roughness did not chang by h at-treatm nt at 1150 °C (J. Mat r. R s. (1990) Vol.5, No.9, p.1918).

T. Sakamoto, et al. report d that, in an Si-MBE apparatus, steps on the surface of Si waf r decreased, and the luminance at the mirror reflection point in r flective high-speed lectron diffraction increased monotonously with annealing time when

the spontaneously formed oxid film was removed, an Si layer was grown, and the lay r was ann aled in a high vacuum at a high t mperatur (700°C to 1000°C), as a m thod for flatt ning a waf r at an atomic level (Denki Tsushin Gakkai Gijutsu Hokoku (Technical Bulletin of Electro-communication Society) SSD86-25, (1986) p.47).

N. Nakamura, et al. found that rectangular projections and concavities having height and width of about 1 μ m on a surface of Si were deformed during heat treatment in ultra high vacuum in an MBE apparatus (Journal of Applied Physics, Vol. 68 (1990), p.3038). They reported that it is important, for causing the deformation, to eliminate the spontaneously formed oxide film from the Si surface to provide a surface that is clean in atomic level

Aoyama, et al. reported that surface roughness of an Si substrate changed on exposure to hydrogen-diluted, fluorine having been excited by UV light, and in particular, a (III) Si surface was flattened by 0.1 to 0.2 nm in terms of rms, and they suggested that this change is caused by rate-controlled oxidation of the (III) surface and removal of the resulting oxide by HF (Ohyo Butsuri Gakkai Gakujustsu Koenkai (Meeting for Scientific Lectures of Applied Physical Society (1991), 12a-B-6).

Of the above-described flattening methods, a polishing method is illustrated in Figs. I3A and 13B. where the numeral 11 indicates a substrate; 12, irregularity of the substrate surface; 13, a surface state before the polishing; and 14, a surface state after the polishing. As shown in Fig. I3A, the rough surface of monocrystal is removed a certain amount of the surface layer by polishing to obtain a smoothened and flattened surface as shown in Fig. I3B. In other words, polishing for elimination of roughness removes inevitably a certain amount of the surface layer, and variation in the polishing procedure causes disadvantageously variation of the thickness of the polished Surface layer. Therefore, in the case where the outermost surface or vicinity thereto of a monocrystal is utilized, the removal of the surface layer by pollshing is not suitable for smoothening and flattening of the surface insofar as the thickness of the monocrystal is important. Moreover, the surface layer of monocrystal provided by polishing usually has defects such as distortion and dislocation caused by the Polishing operation. After the polishing, such defects need to be eliminated by etching of several hundreds of nm to s veral µm, or at least s v ral tens of nm, of the surfac layer.

The proc ssing of s miconductor devices is practiced in a clean room where contaminating particles ar removed ultimately. The polishing process, which is a dust-forming process, has to be separated from oth r processes, and special car

15

20

25

30

35

has to b taken in delivery of the intermediate products to or from the separated process s. Ther fore th pollshing is not practical in th devic production.

On the other hand, the flattening by heat-treatment in a nitrogen atmosphere involves problems below, because of the treatment in a high temperatur exceeding 1260 °C for a long time:

- (1) A specially designed treating-furnace is required which employs a high-temperature-resistant construction material such as SiC in the semiconductor process, since the temperature exceeds the maximum resistance temperature of a silica tube (a silica tube may be distorted at a temperature of I200 ° C).
- (2) A number of slip lines may be introduced to the monocrystal owing to temperature distribution in the substrate or temperature variation on putting the substrate into or out from the furnace, since the treating temperature is near the melting point of silicon.
- (3) Impurity may diffuse and redistribute, in the case where an impurity such as boron and phosphorus is preliminarily introduced locally in monocrystal layer.

To solve the above problems, a method is being sought which is capable of forming a surface having the same level of flatness as or a higher level than that achieved by polishing, at a temperature for a production process of devices, integrated circuits, etc. or a lower temperature without removal of monocrystal surface.

One of the flattening methods, in which the temperature is adaptable to the process and the surface layer of the monocrystal is not removed, is a heat-treatment in the aforementioned MBE (molecular beam epitaxy) apparatus. This technique, however, relates to a flattening of steps at an atomic layer level (less than 1 nm) or deformation of grooves on a surface by migration of surfac atomic layer, but does not relates to elimination of roughness of about le3 nm to about I nm in terms of p-V. Further, in this technique, a high vacuum or ultra-high vacuum (lower than 1 x 10-11 Torr) is required since the removal of an oxide film on the surface and prevention of re-oxidation are indispensable. However, heating in high vacuum involve problems of nonuniform temperature distribution and long time required for cooling becaus of lack of heat transfer by convention. If heating is conducted with an exothermic body, a further problem of diffusion of an impurity from th exothermic body aris s.

Furthermor, the MBE process is exp nsive in comparison with the polishing tr atm nt because of the necessity of high performance of a pump for the a large volume of an ultra-high vacuum chamber and the cost of the MBE chamber, and addi-

tionally involves the above-mentioned problem of th hat source. Therefore, the MBE is not suitable for industrial mass-production proc sses wh r a large ar a of substrate is treated or a large number substrates are treated simultaneously. Nakamura, et al. (Ohyo Butsuri Gakkai Gakujutsu Koenkai (Meeting for Scientific Lectures of Applied Physical Society (1990), 27a-T-2) pointed out that, in the MBE process, defects are introduced in a high density into monocrystal with deformation over 800°C, which are caused by oxygen and carbon. Thus, at a high temperature where surface atoms can readily move, defects is disadvantageously introduced.

The flattening by hydrogen-diluted fluorine gas which has been excited by UV light leads to improvement by only from about 0.1 to 0.2 nm in terms of rms, and is limited in capability of flattening the roughness, being not capable of improving roughness at the same level as polishing. Furthermore, this method involves plane orientation dependance, and the (100) plane is roughened adversely.

On the other hand, formation of monocrystalline Si semiconductor layer on an insulator is well known as, silicon-on-insulator (SOI) technique. Many investigations have been made thereof since the devices made by the SOI technique have many advantages which are not achievable with a bulk Si substrate for usual Si integrated circuits. The advantages brought about by the SOI technique are as below:

- 1. Ease of dielectric separation, and practicability of high integration,
- 2. High resistance against radioactive rays,
- 3. Low floating capacity, and practicability of high speed operation,
- 4. Practicability of omission of a welling step.
- 5. Practicability of prevention of latching-up,
- Practicability of thin film formation for complete depletion type field effect transistor, and so forth.

In order to realize the aforementioned advantages in device characteristics, investigations have been made on the methods of forming the SOI structure for the last few decades. The results of the investigations are collected in the literature, for example: Special Issue: "Single-crystal silicon on non-single-crystal insulators"; edited by G.W. Cullen, Journal of Crystal Growth, Vol. 63, No. 3, pp. 429-590 (1983).

Form rly, the SOS techniqu (silicon-on-sapphire) is known which hetero pitaxially forms Si on monocrystalline sapphire substrate by CVD (ch mical vapor d position). This technique, although successful as completed SOI technique, is not widely applied because of many crystal defects caused by insufficient coh rency of the lattice at

10

15

20

25

30

35

45

th interface between the Si layer and the underlying sapphire substrate, migration of aluminum from the sapphir substrate to th Si layer, and, abov all, th high cost of the substrat and difficulty in enlarging the area thereof.

In recent years, the SOI structure without use of the sapphire substrate is going to be realized. This attempt is made in two methods:

- (1) method of oxidizing a surface of an monocrystalline Si substrate, forming an aperture in the oxidized layer to uncover locally the Si substrate, and growing Si epitaxially in a lateral direction using the uncovered Si as the seed to form an monocrystalline Si layer on the SiO_2 (deposition of Si on SiO_2), and
- (2) method of forming SiO₂ under the monocrystalline Si substrate by use of the monocrystalline Si substrate itself as the active layer (no deposition of Si layer).

The means for practicing the method (I) above include direct epitaxial growth of monocrystalline Si layer in a lateral direction by CVD; deposition of amorphous Si and subsequent epitaxial growth in solid in a lateral direction by heat treatment; growth of amorphous or polycrystalline Si into monocrystalline layer by melting-recrystallization by focusing thereon an energy beam such as electron beam and laser light; and zone melting recrystallization. These methods have both advantages and disadvantages, still involving problems in controllability, productivity, uniformity, and quality of the products. Therefore, none of the above methods has been practiced industrially.

For example, the CVD process requires sacrificial oxidation. The solid growth results in low crystallinity. The beam annealing process involves problems in treating time, control of superposition of th beam and focus adjustment. Among the above methods, zone melting recrystallization is most highly developed, and has been employed for experimental production of relatively large integrated circuits. This method, however, still causes crystal defects in subgrain boundary, etc., and does not giv a minority carrier device.

The method (2) above, which do not use the Si substrate as the seeds for epitaxial growth, are practiced in the three ways below:

1. A surface of a monocrystalline Si substrate is etched anisotropically to form V-shaped grooves on the surface. An oxide film is formed thereon. On the oxide film, a polycrystalline Si layer is deposited in a thickn ss that is nearly the same as the Si substrate. Then the Si substrate is abrad d at the backside to form diel ctrically separated monocrystalline Si r gions surrounded by the V-shaped grooves on the thick polycrystalline Si layer. Although this technique gives satisfactory crystallinity, it involves prob-

I ms in control and productivity in the process of depositing polycrystalline SI in a thickness of as much as s veral hundr d μm and in the process of abrading th monocrystallin SI substrate from the backside to leave only the separate active Si layer.

- 2. An SiO₂ layer is formed on a monocrystalline SI layer by oxygen ion Implantation. This method is called SIMOX (separation by ion implanted oxygen). This process exhibits excellent coherency with the Si process, and is the most highly developed technique at the moment. However, the process requires implantation of oxygen ions as much as 10¹⁸ ions/cm² to form the SiO₂ layer, and the implantation takes long time, so that the productivity is not high, and the wafer cost is high. Furthermore, the product has many remaining crystal defects, and does not have satisfactory quality for industrial production of minority carrier devices.
- 3. The SOI structure is formed by dielectric separation by oxidation of porous Si. In this method, on a surface of a P-type monocrystalline Si substrate, an N-type Si layer is formed in an island shape by proton ion implantation (Imai, et at.: J. Crystal Growth, vol. 63, 547, (1983)), or by epitaxial growth and patterning, then the Ptype Si substrate only is made porous by anodization in an HF solution so as to surround the island-shaped Si regions, and the N-type Si layers are separated dielectrically by accelerated oxidation. This method has disadvantage that the freedom in device design is frequently restricted since the Si region to be separated has to be decided prior to the device production process.

On the other hand, a light-transmissive substrate typified by glass allow Si to grow only into an amorphous or polycrystalline layer under the influence of the disorderness of the crystal structure of the glass, and is unsuitable for production of devices of high performance. Simple deposition of Si on such a substrate will not give excellent monocrystal layer because of the amorphous structure of the substrate. The light-transmissive substrate is important in constructing a light-receiving element for a contact sensor, a projection type liquid crystal image displaying apparatus, and the like. In order to provide a sensor or an image element (picture element) of a display apparatus in higher density. higher resolution, and higher fineness, extremely high performance of th driving devic is required. Therefore, the devic on a light-transmissiv substrat have to b made from monocrystal lay r having exc II nt crystallinity.

In other words, amorphous Si or polycrystalline Si will not gen rally give a driving device which xhibits the satisfactory performance required or to

20

30

35

40

50

be r quired in the future because of many defects in the crystal structure.

On a light-transmissiv substrat, however, none of the abov methods for a monocrystalline Si substrate is suitable for forming a monocrystal layer with good quality.

SUMMARY OF THE INVENTION

An object of the present invention is to provide a process for producing a semiconductor substrate in which surface roughness is eliminated without decreasing the surface layer thickness to an extent achievable by polishing or a higher extent in a method other than usually employed polishing m thods.

Another object of the present invention is to provide a process for producing a semiconductor substrate which has the advantage of the conventional SOI structure and is applicable to various high-performance electronic devices.

Still another object of the present invention is to provide a process for producing a semiconductor substrate which is useful in place of expensive SOS and SIMOX in production of large scale integrated circuits of SOI structure.

A further object of the present invention is to provide a process for producing a semiconductor substrate having, on an insulating layer, an Si layer of crystallinity and surface flatness as high as monocrystalline wafer with prominence in productivity, uniformity, controllability, and cost.

A still further object of the present invention is to provide a process for producing a semiconductor substrate having, on a transparent substrate (light-transmissive substrate), an Si layer of crystallinity as high as monocrystalline wafer with prominence in productivity, uniformity, controllability, and cost.

A still further object of the present invention is to provide a process for producing a semiconductor substrate, comprising heat-treating a substrate having a monocrystal thereon at a temperature lower than the melting point of the monocrystal in a reducing atmosphere, and a process for producing an SOI semiconductor substrate by utilizing the above-mentioned process.

BRIEF DESCRIPTION OF THE DRAWINGS

Figs. 1A to 9C are schematic views illustrating respectiv ly an example of the process for producing the similar micronductor substrate of the present invention.

Figs. 10 and II ar respectiv ly a graph showing, the properties of porous silicon in etching.

Fig. 12 is a graph showing the change of por density by heat-treatment.

Figs. 13A and 13B illustrate schematically flattening of the surface of a semiconductor substrate by conventional pollshing proc ss.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

A first embodiment of the process for producing a semiconductor substrate of the present invention is a process comprising heat-treating a substrate having a monocrystal thereon in a reducing atmosphere to flatten and smooth the surface of the monocrystal.

The present invention involves a process for producing a semiconductor substrate having SOI structure by using the first embodiment of the present invention.

A second embodiment of the process for producing a semiconductor substrate of the present invention is a process comprising the steps of porousifying a silicon base member, forming a non-porous monocrystalline silicon layer on the porous material to prepare a first substrate, heat-treating the first substrate at a temperature lower than the melting point of the nonporous monocrystalline silicon in a reducing atmosphere, bonding the surface of the nonporous monocrystalline silicon layer onto a second substrate with interposition of an insulating layer, and then removing the porous silicon by etching.

A third embodiment of the process for producing. a semiconductor substrate of the present invention is a process comprising the steps of porousifying a silicon base member, forming a non-porous monocrystalline silicon layer on the porous material to prepare a first substrate, bonding the surface of the nonporous monocrystalline silicon onto a second substrate with interposition of an insulating layer, removing the porous silicon by etching, and heat-treating the bonded substrates at a temperature lower than the melting point of the nonporous monocrystalline silicon in a reducing atmosphere.

A fourth embodiment of the process for producing a semiconductor substrate of the present invention is a process comprising the steps of porousifying a silicon substrate, heat-treating the porousified silicon substrate at a temperature lower than the melting point thereof in a reducing atmosphere to change the surface layer of the porousified silicon substrate into a nonporous monocrystalline silicon layer, bonding the nonporous monocrystalline silicon layer onto the surface comprising an insulating material of a second substrate, and removing the porousified portion of the silicon substrate by etching.

According to the first embodiment of the present invention, a surface of a monocrystal is

40

50

made as flat as the surface of a monocrystalline waf r with substantially no chang of th thickness of the monocrystal without introducing a processing d formation lay r and without introducing crystal defects such as slip lines, because the crystal removal procedure like as polishing is not employed. The roughness of several tens of nm in terms of the p-v value can be made flat, which is not achievable by heat-treatment in ultra-high vacuum r by treatment with a UV-excited hydrogendiluted fluorine gas. This process, which utilizes heat treatment, is capable of treating simultaneously a number of substrates. Further, since the treating t mperature is similar to that employed in usual semiconductor process and no ultra-high vacuum is necessary, this process can be realized with an ordinary heat-treating apparatus in a semiconductor process line without additional equipment investment. The heat-treatment can be practiced in continuation to other processes. According to this invention, the roughness in a local monocrystal region on a surface of a substrate which cannot be made flat by polishing can be made flat.

The second to fourth embodiments enable production of a substrate having SOI structure with retention of the advantages of the first embodiment.

According to the second embodiment of the present invention, the surface of the first substrate to be bonded to the second substrate is made as flat as the surface of a wafer by heat treatment in a reducing atmosphere, whereby the nonuniformity of th bonding is avoided and the strong and uniform bonding is realized throughout the whole substrate.

According to the third embodiment of the pr sent invention, the surface after removal of porous silicon layer by etching can he made as flat as the surface of a wafer by heat treatment in a reducing atmosphere.

According to the fourth embodiment of the present invention, a silicon substrate is made porous and then is heat-treated at a temperature lower than the melting point thereof to transform th surface of the porous silicon substrate into a nonporous monocrystalline silicon layer, whereby a monocrystalline silicon layer can be formed with high crystallinity on a porous silicon substrate, without using a source gas such as silane.

The present invention is described below in mor detail by reference to specific embodiments and xamples.

Embodiment I

The inventors of the pres nt invention inv stigated h at tr atm nt for elimination of fine roughness of a surface of monocrystalline silicon, and have found that the roughness of th surface of

monocrystallin silicon can be liminated by heat treatment in a reducing, atmospher at a t mperature not higher than that in a d vice process without application fultra-high vacuum. Th "reducing atmosphere" herein includes a hydrogen-containing atmosphere, and a hydrogen atmosphere, but is not limited thereto. As the results of detailed observation of changes of the surface roughness by high-resolution scanning electron microscopy and interatomic force microscopy, it was found that the surface roughness before the heat treatment as shown in Fig. IA is decreased by heat treatment in a reducing atmosphere, and a thin monocrystal layer having a flat surface is obtained as shown in Fig. 1B. In Figs. IA and IB, the numeral 101 denotes a substrate; 102, roughness on the surface of the substrate, 103, a surface state before the heat treatment; and 104, a surface state after the heat treatment. In the case where the surface roughness is eliminated by polishing, the thickness of the monocrystal layer may vary in the plane. On the contrary, in the heat treatment in a reducing atmosphere of the present invention, the fine roughness only is removed, but the thickness of the monocrystal itself does not vary. Accordingly, the surface flattening by the heat treatment does not cause additional thickness variation.

It was found that, the surface roughness of monocrystalline silicon having a roughness of about several nm to several tens of nm in terms of the p-v value in a cycle of several nm to several hundreds of nm (Fig. 1A) can be flattened by heat treatment in a reducing atmosphere to a p-v value of several nm or less, or 2 nm or less under optimum conditions (Fig. 1B) as shown by the numeral 104 in Fig. 1B, which is the same flatness level as monocrystalline silicon wafer. This phenomenon is considered to be reconstruction of the surface rather than etching. On a rough surface, a number of edge-shaped portions exist which have high surface energy, and faces of higher order of orientation than the plane orientation of the crystal layer are exposed in great numbers on the surface. The surface energy in such regions Is higher than the surface energy depending on the plane orientation of the surface of monocrystal. In the heat treatment in a reducing atmosphere, a spontaneously formed oxide film on the surface is removed by reducing action of the reducing substance such as hydrogen and does not deposit again during the neat treatment. It is considered that the energy barri r to th movement of surfac Si atom is lowered thereby, which causes movement of Si atoms excit d by the thermal en rgy to construct flat surface having lower surface energy. Therefore, the flattening of the present invention is promoted at the lower index of the plane orientation of the monocrystal surface.

25

Consequ ntly, the surface is flatt n d sufficiently at a temp rature of 1200 °C or lower wher the surface cannot be flatt ned in a nitrog n atmospher or a rar gas atmosphere. The temperature of the heat treatment for flattening the surface in the present invention depends on the gas composition, the pressure, etc., and is usually in the rang of from 300 °C to the melting point of the flattened substance, preferably not lower than 500 °C and not higher than I200 °C. With a higher reducing ability of the atmosphere, the flattening can he promoted at a higher pressure. The pressure is usually not higher than the normal atm spheric pressure, more preferably not higher than 200 Torr. When the flattening proceeds at a low rate undesirably, the heat treatment time is prolonged to obtain a desired flatness of the surfac .

This phenomenon is initiated by heat treatment on a clean surface. If the surface is covered with a thick spontaneous oxide film, the initiation of surface flattening is hastened by preliminary removal of the oxide film, e.g., by etching with dilute hydrofluoric acid prior to the heat treatment.

The resulting flat surface of monocrystal is, suitable for use for production of semiconductor elements.

Embodiment 2

A method is explained in which an Si substrate is made porous and thereon a monocrystal layer is grown epitaxially.

A monocrystalline Si substrate 201 is made porous entirely as shown in Fig. 2A or partially as shown in Fig. 4A by anodization in an HF solution. While the density of monocrystalline Si is 2.33 g/cm³, the density of porous monocrystalline Si lay r can be varied in the range of from 0.6 to 1.1 g/cm³ by varying the concentration of the HF solution in the range of from 50 to 20 %. The porous monocrystalline Si layer tends to be formed readily in a P-type substrate by the reasons below. The por s formed have an average diameter of from about 50 Å to about 600 Å according to observation by transmission electron microscopy.

The porous monocrystalline Si was found in the course of study on electropolishing of semiconductors in the year 1956 by A. Uhlir et al. (A. Uhlir t al.: Bell Syst. Tech. J., vol. 35, p. 333 (1956)). The dissolving reaction of Si in anodization was studied by Unagami, who reported that positive holes are required for anodic reaction of Si in an HF solution and the reaction proceeds according to the formulas below (T. Unagami: J. El ctrochem. Soc., vol. 127, p. 476 (1980):

$$Si + 2HF + (2-n)e^+ \rightarrow SiF_2 + 2H^+ + ne^-$$

$$SiF_2 + 2HF \rightarrow SiF_4 + H_2$$

 $SiF_4 + 2HF \rightarrow H_2SiF_6$

or oth rwise,

Si + 4HF +
$$(4-\lambda)e^+ \rightarrow SiF_4 + 4H^+ + \lambda e^-$$

SiF₄ + 2HF \rightarrow H₂SiF₅

where e^+ and e^- represent respectively a positive hole and an electron; n and λ represent respectively the number of the positive holes required for dissolving one atom of monocrystalline Si, and it was also reported that porous monocrystalline Si is formed if n > 2, or $\lambda > 4$.

Accordingly, a P-type Si having positive holes is considered to be more readily made porous. This selectivity in porous structure formation has already been actually proved by Nagano et al. (Nagano, Nakajima, Yasuno, Ohnaka, and Kajihara: Densi Tsushin Gakkai Gijutsu Kenkyu Hokoku (Technical Builetin of Electronic Communication Society) vol. 79, SSD 79-9549 (1979)) and K.Imai: Solid State Electronics, vol. 24, p. 159 (1981)). Thus, P-type silicon having positive holes can be selectively made porous.

On the other hand, it was also reported that high-concentration N-type monocrystalline Si can also be made porous (R.P. Holmstorm, I.J.Y. Chi: Appl; Phys. Lett. vol. 42, p. 386 (1983)). Therefore it is important to select the substrate capable of being made porous, regardless of whether the structure is of P-type or of N-type.

Subsequently, a thin monocrystal layer 202 is formed on the porous substrate surface by epitaxial growth according to various growth method.

In the porous Si layer, pores of about 600 Å in average diameter are formed according to observation with transmission electron microscopy. Although the density thereof is half or less than the density of monocrystalline Si, the monocrystallinity is retained, and epitaxial growth of monocrystalline Si is practicable on the surface. However, rearrangement of the internal pores occurs at a temperature above 1000 °C, which impairs the characteristics of enhanced etching. Therefore, the epitaxial growth of monocrystalline Si is preferably conducted by a low temperature growth process such as molecular beam epitaxial growth, plasma CVD, thermal CVD, photo-assisted CVD, bias sputtering, liquid crystal growth, etc.

In the epitaxial growths on the porous Si, the strain which arises in the het roepitaxial growth can b relax d, and d velopm nt of d f cts can be pr vented by the structural feature of th porous Si.

The thin monocrystalline layer which is grown pitaxially on the porous Si, may have a rough surface that is not suitable for the bonding to the

50

second substrat mention d later, or may locally be pe lled off in the later h at treatm nt or tching process even if it has been once bond d to ach other, dep nding on the shap of the porous Si, as shown in Fig. 2A and Fig. 4A.

In the present invention, after the thin monocrystal layer is formed on the porous layer, the substrate having the thin monocrystal layer formed thereon is heat-treated in a reducing atmosphere to flatten the surface of the thin monocrystalline Si layer as shown in Fig. 2B and Fig. 4B. The conditions of the heat treatment are the same as in Emb diment 1.

In Fig. 2A and Fig. 2B, the numeral 205 denotes porous silicon; 202, epitaxially grown silicon; 206 the surface state of the epitaxially grown silicon; and 207, the surface state after heat treatment of the pitaxially grown silicon.

The surface of the monocrystalline Si layer on the porous Si substrate is bonded to another substrate 203 such as a silicon substrate having an insulating layer on the surface thereof, a lighttransmissive substrate typified by glass, and the lik as shown in Fig. 2C and Fig 4C. Prior to the bonding, the oxide layer may be formed on the surfac of the monocrystalline layer on the porous Si as the interface between the monocrystalline silicon layer and the insulating layer. This oxide lay r will play an important role in production of a device. That is, the level of underlying interface formed by oxidation of the monocrystalline silicon lay r of the active Si layer can be made lower than the level of the bonding interface, in particular, of the int rface with glass. Thus by locating the bonding interface apart from the active layer, the level which may be generated at the bonding interface can be kept apart, thereby the performance of the resulting electron device is remarkably improved. An oxide layer formed on the surface of the monocrystalline Si layer on the porous Si may be bonded to a desired substrate such as an Si substrate.

Subsequently, the porous Si substrate 15 is entirely removed by chemical etching. The thin monocrystalline Si layer remains on an insulating layer on the substrate or a light transmissive substrate as shown in Fig. 2D. Prior to the etching, an etching preventing film may be formed, if necessary. For example, the bonded two substrates are entir ly coated by deposition of Si₃N₄, and then th Si₃N₄ layer on the surface of the porous Si substrate is removed. Apiezon Wax as anoth r tching pr venting film may b us d in plac of th Si₃N₄ lay r.

The porous lay r has a density of half or I ss than the d nsity of nonporous layer because of the great numb r of voids formed in the interior thereof, and has a greater surface area for the v lum.

Cons qu ntly, the rate of the chemical tching is significantly greater than that of a normal non-porous monocrystallin lay r.

The known me thods for the etching of porous SI include:

- (I) Etching of porous Si with an aqueous NaOH solution (G.Bonchil, R.Herino, K.Barla, and J.C.Pfister: J. Electrochem. Soc., vol. 130, no. 7, p. 1611 (1983)), and
- (2) Etching of porous Si with an etching solution capable of etching monocrystalline Si.

In the above method (2), a hydrofluoric nitric acid type etching solution is used normally. With this etching solution, the etching proceeds through oxidation of Si by nitric acid to form SiO_2 and subsequent etching of the resulting SiO_2 with hydrofluoric acid as shown below:

Si + 20
$$\rightarrow$$
 SiO₂
SiO₂ + 4HF \rightarrow SiF₄ + 2H₂O

The known methods for etching crystalline Si include etching with an ethylenediamine type, KOH type, or hydrazine type of etching solution as well as the above hydrofluoric nitric acid type etching solution.

The selective etching of porous Si, which is particularly effective and important in the present invention, employs hydrofluoric acid or a buffered hydrofluoric acid which does not have etching action for crystalline Si. In this etching, an aqueous hydrogen peroxide solution may be used additionally as an oxidizing agent. The reaction rate can be controlled by changing the ratio of addition of the hydrogen peroxide. An alcohol may be added which serves as a surface active agent to remove instantaneously the bubbles of the gaseous reaction product from the etching surface, and enables selective etching of porous Si uniformly and efficiently.

In the case where the Si substrate 401 is changed partially to porous Si 405 as shown in Figs. 4A to 4C, the nonporous part of the Si is removed at the backside of the substrate opposite to the porous layer to uncover the porous layer 405 by grinding or polishing, or etching with a mixture of hydrofluoric acid, nitric acid and acetic acid, or combination thereof as usually employed in Si wafer Production (Fig. 4D), and then the porous Si layer is removed by chemical etching to leave the thin monocrystalline Si layer on the surface of the insulating layer on a substrate or on the surface of a light-transmissive substrate as shown in Fig. 4E. In Figs. 4A to 4E, th numeral 402 denotes epitaxially grown Si; 406, a surfac stat of epitaxially grown Si; and 407, a surface state after the heat treatment of the epitaxially grown Si.

20

35

45

Fig. 2D and Fig. 4E illustrat r spectively a s miconductor substrate according to the pr sent invention, in which a monocrystallin lay r 202, 402 having crystallinity similar to a silicon waf r is formed in a shape of uniform flat thin layer in a large area on the substrate having an insulating lay r on the surface 203, 403 or a light-transmissiv substrate 203, 403 over the entire wafer.

The resulting semiconductor substrate is useful for production of dielectrically isolated electronic el ments.

Embodiment 3

Firstly, as shown in Fig. 3A, a low-impurity-concentration layer 302 is formed epitaxially by a thin film growth method, or otherwise an N-type monocrystalline layer 302 is formed by proton ion implantation, on the surface of a P-type monocrystalline Si substrate 301.

Then as shown in Fig. 3B, the P-type monocrystalline Si substrate 301 is made porous from the backside by anodization by use of an HF solution to form a porous Si layer 303. While the density of monocrystalline Si is 2.33 g/cm³, the density of porous monocrystalline Si layer 15 can be varied in the range of from I.I to 0.6 g/cm³ by varying the concentration of the HF solution in the range, of from 50 to 20 %. The porous layer is formed in a P-type substrate as described before.

Thereafter, the surface roughness is lessened by heat treatment of the nonporous Si monocrystal-line Si layer in a reducing atmosphere in the same manner as in the above Embodiment 2 (Fig. 3C), and the surface of the monocrystalline Si layer on the porous Si substrate or an oxidized surface of the monocrystalline Si layer is bonded to another substrate 304 having insulating layer on the surface ther of as shown in Fig. 3D. Otherwise, an oxidized surface formed on the monocrystalline Si layer surface is bonded to any desired substrate such as an Si substrate.

The porous layer of the porous Si substrate 303 is removed entirely by etching to obtain a substrate having an insulting layer thereon and thin monocrystalline silicon layer further thereon as shown in Fig. 3E.

Fig. 3E illustrates a semiconductor substrate, in which a monocrystalline layer 302 having crystallinity similar to a silicon wafer is formed in a shape of uniform flat thin layer in a large area on a substrate 304 having an insulating layer on the surface or a light-transmissiv substrate 304 over the ntir wafer.

The r sulting s miconductor substrate is us ful for production of dielectrically isolated electronic elements.

In the abov mbodiments, an N-type lay r is formed befor the porous structure formation, and subsequently the P-type portion of the substrationly is a lective ly made porous.

Embodiment 4-I

A monocrystalline Si substrate 701 is made porous entirely as shown in Fig. 5A or partially as shown in Fig. 7A by anodization in an HF solution in the same manner as in above Embodiments to form a porous layer 505, 705.

Subsequently, a thin monocrystal layer 502, 702, is formed on the porous substrate surface by epitaxial growth according to any growth method.

Known methods for the etching of porous Si, and the effective method for selective etching of the porous Si in the present invention are described before.

Subsequently, as shown in Fig. 5B and Fig. 7B, the surface of the monocrystalline Si layer is bonded to the surface of another substrate 503, 703, for example, a silicon substrate as a base material having an insulating layer on the surface, or a light-transmissive insulating substrate typified by glass.

Prior to the bonding, an oxide layer may be formed on the surface of the monocrystalline layer on the porous Si as the interface between the monocrystalline silicon layer and the insulating layer. This oxide layer will play an important role in production of a device. That is, the level of the underlying interface formed by oxidation of the monocrystalline silicon layer of the active Si layer can be made lower than the level of the bonding interface, in particular, of the interface with glass. Thus by locating the bonding interface apart from the active layer, the energy level which may be generated at the bonding interface can be kept apart, thereby the performance of the resulting electron device is remarkably improved. The oxide layer formed on the surface of the monocrystalline Si layer on the porous Si may be bonded to a desired substrate such as an Si substrate.

Subsequently, the porous Si substrate 505 is removed by chemical etching. The thin monocrystalline Si layer remains on an insulating layer on the substrate or a light transmissive substrate as shown in Fig. 5C. Prior to the etching, an etching preventing film may be formed, if necessary. For example, the bonded two substrates are entirely coated by deposition of Si₃N₄, and then the Si₂N₄ lay r on the surface of th porous Si substrate is r moved Apiezon Wax may b used in place of th Si₃N₄.

In th cas wher th Si substrate is partially made porous as in the steps shown in Figs. 7A and 7B, the nonporous part of the Si is removed at the, backside of the substrate opposite to the porous

25

40

50

layer to uncover the porous layer by grinding or polishing, or tching with a mixtur of hydrofluoric acid, nitric acid and acetic acid, or combination thereof as usually employed in Si waf r production, and then the porous Si layer is removed by chemical etching to leave the thin monocrystalline Si lay r on the surface of the insulating layer on a substrate or a light-transmissive substrate as sh wn in Fig. 7C.

The resulting substrate which has a nonporous monocrystalline silicon layer on an insulating layer after th removal of the porous silicon layer is heat-treat d in a reducing atmosphere to form a monocrystalline silicon layer having a flat surface on a substrate having an insulation layer on the surface or on a light-transmissive substrate as shown in Fig. 5C and Fig. 7C.

The inventors of the present invention investigated the heat treatment for eliminating the minute roughness of the surface of thin nonporous monocrystalline silicon, and found that the roughness of th nonporous monocrystalline silicon surface can be eliminated by heat treatment in a reducing atmosphere at a temperature usually employed in device processes or lower. The reducing atmosph r herein includes a hydrogen-containing atmosphere and a hydrogen atmosphere, but it is not limited thereto. As the result of the detailed observation of the state of the surface roughness in heat treatment in various atmospheres by high r solution scanning electron microscope, interatomic force microscope, etc., it was found that the irregularity of the surface before the heat treatment is lessened by heat treatment in a reducing atmosphere, and the surface of the thin monocrystal layer becomes flat as shown in Fig. 5C and Fig. 7C. If the surface roughness is eliminated by polishing or a like method, the thickness of the monocrystal layer tends to vary throughout the layer. On the contrary, in the present invention, the minute irregularity of the surface can be eliminated without causing variation of layer thickness by the heat treatment in a reducing atmosphere.

In the microscopic structure of the surface of a nonporous monocrystalline silicon, periodical irregularity of several nm to several tens of nm in height difference in a cycle of from several nm to several hundreds of nm is observed. The heat treatment in a reducing atmosphere gives a flat surface of height difference of not more than several nm, or not more than 2 nm under optimum conditions. This ph nom non is considered to be reconstruction of the surface rather than etching. On a rough surface, a number of edge-shaped portions exist which have high surface energy, and faces of higher order of orientation than the plane orientation of the crystal lay reare exposed in great numbers on the surface. The surface energy in such regions is

higher than the surface nergy at the plane or intation of the surface of the first substrate. In the heat treatment in a reducing atmospher, a spontaneously formed oxide film on the surface is removed by reducing action of the reducing substance such as hydrogen and does not deposit again during the heat treatment. It is considered that the energy barrier to the movement of surface Si atoms is lowered thereby, which causes movement of Si atoms excited by the thermal energy to construct flat surface having lower surface energy.

Consequently, the surface is flattened sufficiently at a temperature of 1200 °C or lower where the surface cannot be flattened in a nitrogen atmosphere or a rare gas atmosphere. The temperature of the heat treatment for flattening the surface in the present invention depends on the gas composition, the pressure, etc., and is usually in the range of from 300 °C to the melting points of the flattened substance, preferably not lower than 500 °C and not higher than 1200 °C. With a higher reducing ability of the atmosphere, the flattening can be promoted at a higher pressure. The pressure is usually not higher than the normal atmospheric pressure, more preferably not higher than 200 Torr.

This phenomenon is initiated by heat treatment on a clean surface. If the surface is covered with a thick spontaneous oxide film, the initiation of surface flattening is hastened by preliminary removal of the oxide film, e.g., by etching with dilute hydrofluoric acid prior to the heat treatment.

Fig. 5D and Fig. 7D illustrate respectively a semiconductor substrate, in which a monocrystal-line Si layer 502, 702 having crystallinity similar to a silicon wafer is formed in a shape of uniform flat thin layer in a large area on the substrate having an insulating layer on the surface 503, 703 or a light-transmissive substrate 503, 703 over the entire wafer.

The resulting semiconductor substrate is useful for production of dielectrically isolated electronic elements.

Embodiment 4-2

Firstly, as shod in Fig. 6A, a low-impurity-density layer 602 is formed epitaxially by a thin film growth method, or otherwise an N-type monocrystalline layer 602 is formed by proton ion implantation, on the surface of a P-type monocrystallin Si substrate 601.

Then as shown in Fig. 6B, the P-type monocrystalline Si substrate 601 is made porous from the backsid by anodization by us of an HF solution to form a porous Si layer 603. While the density of monocrystallin Si is 2.33 g/cm³, the density of porous monocrystallin Si layer 15 can

20

25

35

40

50

55

b varied in the range of from 1.1 to 0.6 g/cm³ by varying the conc ntration of the HF solution in the rang of from 50 to 20 %. The porous lay r is formed in a P-typ substrat as described before.

The surface of the monocrystalline Si layer on the porous Si substrate or an oxidized surface of the monocrystalline Si layer is bonded to another substrate 604 having an insulating layer on the surface thereof, or otherwise, the oxidized surface of the monocrystalline Si layer is bonded to any desired substrate such as an Si substrates shown in Fig. 6C.

The porous layer of the porousified Si substrat 603 is removed entirely by etching to obtain a substrate having an insulting layer thereon and thin monocrystalline silicon layer further thereon as shown in Fig. 6D.

Then in the same manner as in Embodiment 4-1, the substrate having the nonporous monocrystal-line silicon layer on the insulating layer after the removal of the porous silicon is heat-treated in a reducing atmosphere to improve the surface roughness, thereby obtaining a semiconductor substrate of the present invention as shown in Fig.6E. Consequently, a monocrystalline Si layer 602 having crystallinity similar to a silicon wafer is formed in a shape of uniform flat thin layer in a large area on a substrate 604 having an insulating layer on the surface or a light-transmissive substrate 604 over the entire wafer.

The resulting semiconductor substrate is useful for production of dielectrically isolated electronic elements.

In the above embodiments, an N-type layer is formed before the porous structure formation, and subsequently the P-type portion of the substrate only is selectively made porous by anodization.

Embodiment 5

A monocrystalline Si substrate is made porous as shown in Fig. 8A by anodization in an HF solution. In the porous structure formation, the entire, the front surface, or the both surfaces of the substrate may be made porous. Then, the substrate is heat-treated in a non-oxidizing atmosphere or in vacuo at a temperature lower than the melting point of the Si substrate to transform the surface layer of the porous monocrystalline Si substrate 80 into a thin nonporous monocrystal layer 802.

Fig. 10 shows the dependency of the layer thickn ss by tching of th porous Si and that of the monocrystalline Si on tching tim in the case wher th porous Si and th monocrystallin Si are r sp ctiv ly tch d by immersion in a mixture of a buffered hydrofluoric acid, alcohol, and an aqueous hydrogen peroxide solution without stirring.

The porous structure formation and the tching step are xplained specifically.

The porous Si is pr pared by anodization. Th anodization conditions are shown below. The starting material of the porous Si prepared by anodization is not limited to monocrystalline Si, but may be Si of any other crystal structure.

Applied voltage:

2.6 V

Current density:

30 mA+cm⁻²

Anodization solution:

 $HF:H_2O:C_2H_5OH = 1:1:1$

Time:

2.4 hours

Porous Si thickness:

300 µm

Porosity:

56 %

The porous Si prepared under the above conditions was immersed in a 10:6:50 mixture of buffered hydrofluoric acid (4.5% HF + 36% NH₄F + H₂O), alcohol, and aqueous 30% hydrogen peroxide solution at room temperature without stirring. The decrease of the thickness of the porous Si was measured later. The progress of the etching is shown by blank circles (O) in Fig. 10. The porous Si was etched rapidly in the etched layer thickness of 83 μm in 40 minutes and 140 μm in 120 minutes uniformly with retention of satisfactory surface property. The etching rate depends on the concentration of the solution and the temperature.

As described before, the addition of alcohol allows the instantaneous removal of bubbles formed by the etching reaction from the etched surface without stirring, thus enabling uniform and effective etching of the porous Si. The addition of the aqueous hydrogen peroxide solution accelerates the oxidation of silicon, and thereby increases the reaction velocity in comparison with the case where hydrogen peroxide is absent. Further the reaction velocity can be controlled by changing the ratio of the hydrogen peroxide solution.

Separately, nonporous SI of 500 μ m thick was immersed in the 10:6:50 mixture of buffered hydrofluoric acid (4.5% HF + 36% NH₄F + H₂O), alcohol, and aqueous 30% hydrogen peroxide solution at room temperature without stirring. The decrease of the thickness was measured later. The progress of the etching is shown by solid circles (\bullet) in Fig. 10. The nonporous Si was etched by not more than 100 Å ν n after 120 minutes of tching.

Aft r the etching, the porous Si and the non-porous Si w r wash d with water. The surfaces contain d no impurity according to microanalysis by secondary ion mass spectrometry.

The alcohol in the invention includes ethyl alcoh I, and other alcohols such as isopropyl alcohol

20

25

which causes no inc nvenience and exhibits the eff ct of addition of the alcohol.

Th inventors of the present invention observed the change of the structur of the porous lay r during the heat treatment by changing the atmosphere and the like by use of high-resolution scanning electron microscopy and other means. As the results, it was found that the number of pores on the surface of the porous layer decreases by heat treatment in a reducing atmosphere or in vacuo, progress depending on the conditions. For example, as shown in Fig. 12, the pores decreased in number with lapse of time, and finally disapp ared, whereby a thin monocrystal layer having a smooth surface was formed. In the heat treatment f the surface and the portion near the surface of the porous Si substrate formed by anodization, the pores disappears so as to lower the surface energy to flatten the surface, thereby the thin nonporous monocrystal layer being formed.

The nonporous monocrystal layer having a flat surface was confirmed to have the same orientation as the substrate by RHEED and electron beam channeling pattern.

This phenomenon is promoted at a higher temperatur and at a lower pressure. The reducing atmosphere herein includes hydrogen-containing atmosphere and a hydrogen atmosphere, but is not limited thereto. The temperature for the heat treatment depends on the gas composition, the pressure, etc., and is usually in the range of from 300 °C, preferably 500 °C to the melting temperature. With a higher reducing ability of the atmosphere, the flattening can be promoted at a higher pressure, and usually the flattening is conducted at the normal atmospheric pressure or lower, more pr ferably not higher than 200 Torr. There is no lower limit of the pressure. Ultra-high vacuum is not especially necessary. The term "in vacuo" in the present invention means the pressure of 1×10^{-3} Torr or lower, preferably 1×10^{-5} Torr or lower without introducing an atmospheric gas in a state of no leak of the reaction chamber.

This phenomenon is initiated on a clean surface by heat treatment. If the surface is covered with a thick spontaneously formed oxide film, the initiation of the surface flattening is hastened by preliminary elimination of the oxide film, e.g., by etching with hydrofluoric acid prior to the heat tr atm nt.

On the surface of another Si substrate 803, an oxide layer 804 is form d as shown in Fig. 8B. Then the Si substrat 803 having the oxid layer 804 is bond d to the surface of the monocrystallin Si layer 802 on the porous Si substrat 801. The bonding is conducted by bringing the cleaned surfaces into contact with each other and then heating in an oxygen atmosphere or a nitrogen atmosphere.

sphere.

Prior to the bonding, an oxide layer 806 may b formed on the nonporous monocrystalline silicon layer 802, which is then bonded to the Silicon layer 802, which is then bonded to the Silicon layer 806 is formed for the purpose of lowering the interfacial level of the monocrystalline Silicon 802 which is the final active layer. In this case, another Siliconstrate may have an oxide layer thereon, or not.

Prior to the etching, an etching preventing film 805 of Si_3N_4 may be deposited, if necessary, to cover the entire of the two bonded Si substrates, and the Si_3N_4 on the porous layer only is removed. In place of the Si_3N_4 layer, Apiezon Wax may be used as the etching protection film. Later, the porous Si substrate 801 is completely removed by etching, leaving the thin monocrystalline Si layer 802 on the SiO_2 .

Prior to the etching, the porous Si substrate 801 may be made thin by mechanical working such as polishing or grinding from the backside. In particular, when not the entire Si substrate is made porous, the substrate is preferably made thin by mechanical processing so as to expose the porous layer.

Fig. 8C shows a semiconductor substrate provided according to the present invention. After removal of the etching preventing film 805 of Si₃N₄, a monocrystalline Si layer 802 having crystallinity in the same level as of a silicon wafer is formed in a form of flat uniform thin film in a large area over the entire area of the wafer. If necessary, the thin monocrystal layer made thicker by epitaxial growth on the monocrystalline Si layer. The growth may be conducted in any method including CVD, sputtering, liquid-phase growth, and solid-phase growth.

Embodiment 6

A monocrystalline Si substrate is made porous as shown in Fig. 9A. In the porous structure formation, the entire, the front surface, or the both surfaces of the substrate may be made porous. Then, the substrate is heat-treated in a reducing atmosphere or in vacuo at a temperature lower than the melting point of the Si substrate to transform the surface layer of the porous monocrystalline Si substrate 901 into a thin nonporous monocrystal layer 902.

Fig. Il shows the dependency of the thickness by etching if the porous Si and that of the monocrystalline Si on etching tim in the case when the porous Si and the monocrystalline Si are immers define a mixture of hydrofluoric acid, alcohol, and an aqueous hydrogen peroxide solution without stiming.

The porous structure formation and the tching step are explained specifically.

45

50

10

30

35

40

45

50

55

The porous Si is pr pared by anodization of monocrystalline Si. The anodization conditions are shown below. The starting material of th porous Si prepared by anodization is not limited to monocrystalline Si, but may be Si of any other crystal structure.

Applied voltage:

2.6 V

Current density:

30 mA+cm⁻²

Anodization solution:

 $HF:H_2O:C_2H_5OH = 1:1:1$

Time:

2.4 hours

Porous Si thickness:

300 µm

Porosity:

56 %

The porous Si prepared under the above conditions was immersed in a 10:6:50 mixture of 49% hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution at room temperature without stirring. The decrease of the thickness of the porous Si was measured later. The progress of the etching is shown by blank circles (O) in Fig. II. The porous Si was etched rapidly (107 μ m in 40 minutes and 244 μ m in 80 minutes) uniformly with retention of satisfactory surface property. The etching rate depends on the concentration of the solution and the temperature.

Separately, nonporous Si of 500 μ m thick was immersed in the 10:6:50 mixture of 49% hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution at room temperature without stirring. The decrease of the thickness was measured later. The progress of the etching is shown by solid circl s (•) in Fig. 10. The nonporous Si was etched by not more than 100 Å even after 120 minutes of etching.

Subsequently, a light-transmissive glass substrate 903 is bonded to the surface of the monocrystalline Si layer 902 on the porous Si substrate 901. The bonding is conducted by bringing the cleaned surfaces into contact with each other and then heating in an oxygen atmosphere or a nitrogen atmosphere.

Prior to the bonding, an oxide layer 906 may be formed on the nonporous monocrystalline silicon layer 902. The oxide layer 906 is formed for the purpose of lowering the interfacial level of the monocrystalline Si layer 902 which is the final activ lay r.

Prior to the tching, an etching protection film 905 of Si_3N_4 may be d posited, if necessary, to cover the entire Of the two bonded Si substrates, and the Si_3N_4 on th porous lay r only is removed. In place of the Si_3N_4 layer, Api zon Wax may be used as the etching protection film. Lat r, the po-

rous Si substrate 901 is compl tely r moved by etching, leaving the thin monocrystallin Si layer 902 on the light-transmissive glass 903.

Prior to the etching, the porous SI substrate 901 may be made thin by mechanical working such as polishing or grinding from the backside. Particularly when not the entire SI substrate is made porous, the substrate is preferably made thin by mechanical processing so as to expose the porous layer.

The present invention is explained by reference to specific examples without limiting the invention thereto in any way.

Example 1

A 4-inch monocrystalline (100) silicon wafer which has irregularity of about 30 nm in height in a cycle of about 50 nm on the surface observed by interatomic force microscopy was placed in a heat treatment furnace. The furnace was evacuated to a vacuum of 0.01 Torr, and thereto gaseous hydrogen was introduced. The silicon wafer was heattreated at a temperature of 950 °C and a pressure of 80 Torr. The surface was observed by interatomic force microscopy, and the surface roughness of 30 nm before the hydrogen treatment was found to have been improved to 1.5 nm.

The treated Si layer was confirmed to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross section by transmission electron microscopy.

Example 2

A 4-inch monocrystalline (110) silicon wafer which has irregalarity of about 30 nm in height in a cycle of about 50 nm on the surface observed by interatomic force microscopy was placed in a heat treatment furnace. Gaseous hydrogen was introduced into the furnace. The silicon wafer was kept standing for 10 minutes, and then heat-treated at a temperature of II50 °C and a pressure of 760 Torr. The surface was observed by interatomic force microscopy, and the roughness of 30 nm before the hydrogen treatment was found to have been improved to 1.6 nm.

The treated Si layer was confirmed to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross section by transmission electron microscopy.

Example 3

A 4-inch 4*-off monocrystalline (I00) silicon waf r which has irr gularity of about 30 nm in height in a cycle of about 50 nm on the surface observed by interatomic force microscopy was

15

30

40 ,

45

50

55

placed in a heat treatment furnace. The furnace was evacuated to a vacuum of 0.0l Torr, and th reto gaseous hydrogen was introduc d. The silicon waf r was h at-treated at a temperature of 900°C and a pressure of 10 Torr. The surface was observed by interatomic force microscopy and the surfac roughness of 30 nm before the hydrogen treatment was found to have been improved to 1.7 nm.

The treated Si layer was confirmed to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross section by transmission electron microscopy.

Example 4

A 4-inch monocrystalline (III) silicon wafer which has irregularity of about 30 nm in height in a cycle of about 50 nm on the surface observed by interatomic force microscopy was placed in a heat tr atment furnace. The furnace was evacuated to a vacuum of 0.01 Torr, and thereto gaseous hydrog n was introduced. The silicon wafer was heat-treated at a temperature of IIO0 °C and a pressure of 760 Torr. The surface was observed by interatomic force microscopy, and the roughness of 30 nm before the hydrogen treatment was found to have been improved to 1.7 nm.

The treated Si layer was confirmed to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross section by transmission electron microscopy.

Example 5

A 4-inch 4-inch 4*-off monocrystalline (III) silicon wafer which has irregularity of about 30 nm in height in a cycle of about 50 nm on the surface observed by interatomic force microscopy was placed in a heat treatment furnace. The furnace was evacuated to a vacuum of 0.0I Torr, and thereto a gas mixture of nitrogen 90 % and hydrogen 10 % was introduced. The silicon wafer was heattreated at a temperature of 950 °C and a pressure of 50 Torr. The surface was observed by interatomic force microscopy, and the surface roughness of 30 nm before the hydrogen treatment was found to have been improved to 1.9 nm.

The treated Si layer was confirmed to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross section by transmission lectron microscopy.

Example 6

A 4-inch monocrystalline (100) silicon wafer which has irregularity of about 30 nm in height in a cycl of about 50 nm on the surfac observed by

interatomic force microscopy was placed in a heat tr atment furnace. Th r to a gas mixture of nitrogen 90 % and hydrogen 10 % was introduced. The silicon wafer was kept standing for 15 minutes, and the heat-treated at a temperature of 700 °C and a pressure of 0.1 Torr. The surface was observed by interatomic force microscopy, and the surface roughness of 30 nm before the hydrogen treatment was found to have been improved to 1.7 nm.

The treated Si layer was confirmed to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross section by transmission electron microscopy.

Example 7

A sample was provided which had on the surface an exposed monocrystalline (100) silicon area of 100 µm square and was covered in other area with a silicon nitride film. The exposed silicon area had irregularity of about 30 nm in height in a cycle of about 50 nm on the surface observed by interatomic force microscopy. The sample was placed in a heat treatment furnace. Thereto a gas mixture of nitrogen 90 % and hydrogen 10 % was introduced, and the atmosphere in the furnace was completely replaced. The sample was then heattreated at a temperature of 1100 °C and a pressure of 100 Torr. The surface was observed by interatomic force microscopy, and the surface roughness of 30 nm before the hydrogen treatment was found to have been improved to 1.7 nm.

The treated Si layer was confirmed to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross section by transmission electron microscopy.

Example 8

A monocrystalline P-type (100) Si substrate of 600 μ m thick was anodized in 50% HF solution for 20 minutes at a current density of 12 mA/cm². The porous structure was formed at a rate of I.I μ m/min. The P-type (100) Si substrate of 600 μ m thick was made porous to a depth of about 20 μ m.

On the porous P-type (100) Si substrate, an epitaxial Si layer was grown in a thickness of 2 μ m by CVD under the deposition conditions below:

Temperature: 950 • C

Pressure: 80 Torr

Gas: SiH₂ Cl₂/H₂, 0.5/180 (1/min)

Growth rate: 0.33 nm/sec

This Si substrate was h at-tr at d at 950 °C in a hydrog n atmosph re at 80 Torr. The treated sample was valuated for surface flatness by interatomic microscop obs rvation. The surface roughness before th hydrogen treatment of 20 nm was found to be improved to 1.5 nm.

15

20

30

35

40

Th surface of the epitaxial lay r was th rmally oxidized in a depth of 50 nm. On the thermally oxidation film, another monocrystallin silicon substrate was superpos d and the combined substrates were heated in a nitrogen atmosphere at 1000 °C for 2 hours, whereby the two substrates wer bonded tightly.

The substrate having been made porous was polished on the backside to remove the nonporous porti n of the silicon substrate to uncover the por us layer.

The bonded substrates were selectively etched by immersion in a mixture of hydrofluoric acid, alc hol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 20 minutes, the porous Si substrate only was selectively etched and remov d completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the etching solution is as low as about 40 Å or a little less for 20 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the porous Si substrate of $200\mu m$ in thickness was removed, and monocrystalline Si layer of $0.5~\mu m$ thick was formed uniformly with no defect on the silicon oxide surface of the silicon substrate. Unbonded spots were found at a spot density of $0.5~\text{spot/cm}^2$ in comparison with 50 spots/cm² in the case where the heat treatment in hydrogen atmosphere was not conducted.

The resulting Si layer was found to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Example 9

A monocrystalline P-type (100) Si substrate of 200 μ m thick was anodized in 50% HF solution at a current density of 110 mA/cm². The porous structure was formed at a rate of 8.7 μ m/min. The entire of the P-type (100) Si substrate of 200 μ m thick was made porous in 23 minutes.

On the porous P-type (I00) Si substrate, an epitaxial Si layer was grown in a thickness of 0.5 μ m by MBE (molecular beam epitaxy) at a low t mp rature under the d position conditions b low:

Temperature: 700 ° C

Pr ssur : 5×10^{-8} Torr

Growth rate:

0.1 nm/sec

This Si substrate was heat-treated at 950 °C in a hydrogen atmosphere at 80 Torr. The treated sample was evaluated for surface flatness by inter-

atomic microscope obs rvation. The surface roughn ss before the hydrog n tr atm nt of 15 nm was found to be improv d to I.5 nm.

The surfac of the epitaxial layer was th rmally oxidized in a depth of 100 nm. On the thermally oxidation film, another monocrystalline silicon substrate was superposed and the combined substrates were heated in an oxygen atmosphere at 900 °C for 2 hours, whereby the two substrates were bonded tightly.

The bonded two substrates were covered by deposition of Si_3N_4 in a thickness of 0.1 μ m by plasma CVD. The the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 204 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the etching solution is as low as about 40 Å or a little less for 204 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the porous Si substrate of 200 μm was removed, and after removal of the Si₃N₄ layer, monocrystalline Si layer of 0.5 μm thick was formed uniformly with no defect on the silicon oxide surface of the silicon substrate.

The resulting Si layer was found to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Example 10

A monocrystalline P-type (100) Si substrate of 200 μ m thick was anodized in 50% HF solution at a current density of 110 mA/cm². The porous structure was formed at a rate of 8.7 μ m/min. The entire of the P-type (100) Si substrate of 200 μ m thick was made porous in 23 minutes.

On the porous P-type (100) Si substrate, an epitaxial Si layer was grown in a thickness of 5 μ m by plasma CVD at a low temperature under the deposition conditions below:

Gas:

SiH₄

High fr quency power:

120 W

T mperature:

800 · C

50

15

20

30

35

45

50

55

Pr ssure: 1×10^{-2} Torr Growth rate: 2.7 nm/sec

This Si substrate was heat-treated at 1000 °C in a hydrogen atmosphere at 760 Torr. The treated sampl was evaluated for surface flatness by interatomic microscope observation. The surface roughness before the hydrogen treatment of 25 nm was found to be improved to 1.6 nm.

Th surface of the epitaxial layer was thermally oxidized in a depth of 50 nm. On the thermally oxidation film, a fused silica substrate having been optically polished was superposed and the combined substrates were heated in an oxygen atmosphere at 400 °C for 20 hours, whereby the two substrates were bonded tightly.

The bonded substrates were selectively etched by immersion in a mixture of hydrofluoric acid and aqueous hydrogen peroxide solution (1:5) with stirring. In 62 minutes, the porous Si substrate was selectively etched and removed completely with the monocrystalline Si left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by th above etching solution is as low as about 20 Å or a little less for 62 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10^5 or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the porous Si substrate of 200 μm thick was removed, and a monocrystalline Si layer of 0.5 μm thick was formed uniformly with no defect on the fused silica substrate.

The resulting Si layer was found to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Example II

A monocrystalline P-type (I00) Si substrate of 200 μ m thick was anodized in 50% HF solution at a current density of 110 mA/cm². The porous structure was formed at a rate of 8.7 μ m/min. The entire of the P-type (100) Si substrate of 200 μ m thick was made porous in 23 minutes.

On the porous P-type (I00) Si substrate, an pitaxial Si layer was grown in a thickness of I µm by CVD at a low temp rature under the deposition conditions below:

Gas: SiH_4 (0.6 t/min), H_2

1/min)

Temperature: Pressure:

850 ° C 40 Torr

Growth rate:

0.3 μm/min

This Si substrate was h at-treat d at 900 °C in a hydrog n atmosph re at 10 Torr. The treated sample was evaluated for surface flatness by intiratomic microscope observation. The surface roughness before the hydrogen treatment of 18 nm was found to be improved to 1.6 nm.

The surface of the epitaxial layer was thermally oxidized in a depth of 50 nm. On the thermally oxidation film, an optically polished glass substrate having a softening point of about 500°C was superposed and the combined substrates were heated in an oxygen atmosphere at 450°C for 0.5 hour, whereby the two substrates were bonded tightly.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 204 minutes, the porous Si substrate was selectively etched and removed completely with the monocrystalline Si left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the above etching solution is as low as about 40 Å or a little less for 204 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the porous Si substrate of 200 μm thick was removed, and a monocrystalline Si layer of 1 μm thick was formed uniformly with no defect on the glass substrate of a low softening point.

The resulting Si layer was found to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Example 12

A monocrystalline P-type (100) SI substrate of 525 μ m thick was anodized in 50% HF solution for 20 minutes at a current density of 12 mA/cm². The porous structure was formed at a rate of 1.1 μ m/min. The P-type (100) SI substrate of 525 μ m thick was made porous to a depth of 20 μ m.

The partially porous substrate was heat-treated in an oxygen atmosphere at 300 °C for one hour.

On the porous face of the P-type (100) Si substrate, an epitaxial Si layer was grown in a thickn ss of 5.0 μ m at a low temperatur by bias sputtering und r the d position conditions below:

RF fr quency:

100 MHz

High frequency power:

600 W

Temperature:

20

30

300 ° C

Ar gas pressur:

 8×10^{-3} Torr

Growth time:

120 minutes

Target DC bias:

-200 V

Substrate DC bias:

+10 V

This Si substrate was heat-treated at 900 °C in a hydrogen atmosphere at 10 Torr. The treated sample was evaluated for surface flatness by interatomic microscope observation. The surface roughness before the hydrogen treatment of 13 nm was found to be improved to 1.4 nm.

On the surface of the epitaxial layer, was formed a silicon oxide layer of 500 nm thick by thermal oxidation. On the thermally oxidation film, another silicon substrate was superposed and the combined substrates were heated in a nitrogen atmosphere at 1000 °C for 2 hours, whereby the two substrates were bonded tightly.

The substrate having been made porous was polished on the backside to remove the nonporous portion of the silicon substrate to uncover the porous layer.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (I0:6:50) without stirring. In 20 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the etching solution is as low as about 40 Å or a little less for 204 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decrease of the film thickness of the nonporous layer (sev ral tens of Å) by etching was practicably negligible.

Consequently, the 20 μ m-thick porous Si substrat was removed, and a monocrystalline Si layer of 0.75 μ m thick was formed umiformly with no def ct on the 500 nm thick silicon oxide layer on the silicon substrate.

The resulting Si layer was found to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Exampl 13

A monocrystalline P-type (I00) Si substrate of 600 µm thick was anodiz d in 50% HF solution at a current density of 14 mA/cm². The porous structure was formed at a rate of I.3 µm/min. The P-type (100) Si substrate of 600 µm thick was made

porous to a depth of 20 µm.

On the porous face of th P-type (I00) Si substrate, an pitaxial Si lay r was grown in a thickness of I0 μ m at a low temperatur by liquid-phase growth under the growth conditions below:

Solvent:

Sn

Growth temperature:

900 · C

Growth atmosphere:

Ha

Growth time:

20 minutes

This Si substrate was heat-treated at 950 °C in a hydrogen atmosphere at 80 Torr. The treated sample was evaluated for surface flatness by interatomic microscope observation. The surface roughness before the hydrogen treatment of 30 nm was found to be improved to I.8 nm.

On the surface of another monocrystalline silicon substrate, was formed a silicon oxide layer of 1 μ m thick. This silicon substrate at the silicon oxide layer side was superposed on the aforementioned epitaxial Si layer, and the combined substrates were heated in a nitrogen atmosphere at 700 °C for 5 hours, whereby the two substrates were bonded tightly.

The substrate having been made porous was polished on the backside to remove the nonporous portion of the silicon substrate to uncover the porous layer.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (I0:6:50) without stirring. In 20 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the etching solution is as low as about 10 Å or less for 20 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the Si substrate made porous was removed, and a monocrystalline Si layer of 10 μ m thick was formed uniformly with no defect on the silicon substrate.

The resulting Si layer was found to have no crystal d f ct newly introduc d and have satisfactory crystallinity by observation of cross-s ction by transmission el ctron microscopy.

50

20

35

40

45

Example 14

On a P-typ (100) Si substrate of 200 μ m thick, an N-type Si lay r of l μ m thick was formed by proton ion implantation in an H⁺ implantation quantity of 5 \times 10¹⁵ (ions/cm²). This substrate was anodiz d in 50% HF solution at a current density of 100 mA/cm². The porous structure was formed at a rate of 8.4 μ m/min. The entire of the P-type (100) Si substrate of 200 μ m thick was made porous in 24 minutes. In the anodization, the P-type (100) Si substrate only was made porous with the N-type Si lay r unchanged.

This Si substrate was heat-treated at 950 °C in a hydrogen atmosphere at 80 Torr. The treated sample was evaluated for surface flatness by interatomic microscope observation. The surface roughness before the hydrogen treatment of 13 nm was found to be improved to 1.1 nm.

The surface of the N-type monocrystal layer was thermally oxidized in a depth of 50 nm. On the th rmally oxidation film, a fused silica substrate having been optically polished was superposed and the combined substrates were heated in an oxygen atmosphere at 800 °C for 0.5 hour, whereby the two substrates were bonded tightly.

The bonded two substrates were covered by deposition of Si₃N₄ in a thickness of 0.1 µm by low-pr ssure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching. Then the bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 204 minutes, the porous Si substrate was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the etching solution is as low as about 40 Å or a littless for 204 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decr as of the film thickness of the nonporous layer (s veral tens of-Å) by etching was practicably negligible.

Consequently, the porous Si substrate of 200 μ m was removed, and after removal of the Si₂N₄ layer, monocrystalline Si layer of 1.0 μ m thick was form d uniformly with no defect on the silicon oxid surface of the silicon substrate.

Example 15

A monocrystallin P-typ (100) Si substrate of 600 µm thick was anodized in 50% HF solution for 20 minutes at a current density of 5 mA/cm². Th porous structure was formed at a rate of 1 µm/min.

The P-type (100) Si substrat of 600 μm thick was mad porous to a depth of about 20 μm .

On th porous P-type (I00) Si substrat , an epitaxial Si layer was grown in a thickn ss of 2 μ m by CVD under the deposition conditions below:

Temperature:

950 · C

Pressure:

80 Torr

Gas:

SiH₂Cl₂/H₂, 0.5/180 (1/min)

Growth rate:

0.33 nm/sec

The surface of this epitaxial layer was thermally oxidized to a depth of 50 nm. On the thermally oxidation film, another monocrystalline silicon substrate was superposed and the combined substrates were heated in a nitrogen atmosphere at 1000 °C for 2 hours, whereby the two substrates were bonded tightly.

The substrate having been made porous was polished on the backside to remove the nonporous portion of the silicon substrate to uncover the porous layer.

The bonded substrates were selectively etched by immersion in a mixture of hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (l0:6:50) without stirring. In 20 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the etching solution is as low as about 40 Å or a little less for 204 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the porous Si substrate of 20 μm thick was removed, and monocrystalline Si layer of 0.5 μm thick was formed on the silicon oxide surface on the silicon substrate after removal of the Si₃N₄ layer.

This substrate was heat-treated at 950 °C in a hydrogen atmosphere at 80 Torr. The treated sample was evaluated for surface flatness by interatomic microscope observation. The surface roughness before the hydrogen treatment of 20 nm was found to have been improved to 1.5 nm.

The resulting Si layer was found to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Exampl 16

A monocrystallin p-typ (l00) Si substrat of 200 μ m thick was anodiz d in 50% HF solution at a current d nsity of 100 mA/cm². The porous structure was form d at a rat of 8.4 μ m/min. The entire of the P-type (l00) Si substrate of 200 μ m thick

10

20

30

35

40

45

50

55

was made porous in 24 minutes.

The porous substrat was h at-treated in an oxygen atmosphere at 300 °C for one hour.

On the porous P-type (100) Si substrat, an pitaxial Si layer was grown in a thickness of 0.5 μ m by MBE (molecule beam epitaxy) at a low t mperature under the deposition conditions below:

T mperature:

700 ° C

Pressure:

1 x 10⁻⁹ Torr

Growth rate:

0.1 nm/sec

The surface of the epitaxial layer was thermally oxidized to a depth of 100 nm. On the thermally oxidation film, another monocrystalline silicon substrate was superposed and the combined substrat s were heated in an oxygen atmosphere at 900 °C for 2 hours, whereby the two substrates wer bonded tightly.

The bonded two substrates were covered by deposition of Si_3N_4 in a thickness of 0.1 μ m by plasma CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (l0:6:50) without stirring. In 204 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous monocrystalline Si layer by the etching solution is as low as about 40 Å or a little less for 204 minutes. Ther fore, the relative rate of selective etching of the porous layer to that of nonporous layer of 10⁵ or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the porous Si substrate of 200 μ m was removed, and after removal of the Si₂N₄ layer, monocrystalline Si layer of 0.5 μ m thick was formed uniformly with no defect on the silicon oxid surface of the silicon substrate.

This silicon substrate was heat-treated at 950 °C in a hydrogen atmosphere at 80 Torr. The treat d sample was evaluated for surface flatness by interatomic microscope observation. The surface roughness before the hydrogen treatment of 20 nm was improved to 1.5 nm.

Embodiment 17

A monocrystalline P-typ (100) Si substrat of 200 μm thick was anodized in 50% HF solution at a current density of 100 mA/cm². The porous structure was form d at a rate of 8.4 μm/min. The entire of the P-type (100) Si substrat of 200 μm thick was made porous in 24 minutes.

On the porous P-typ (100) Si substrate, an epitaxial Si lay r was grown in a thickness of 5 μ m by plasma CVD at a low temperature under the disposition conditions below:

Gas:

SiH₄

High frequency power:

100 W

Temperature:

800 · C

Pressure:

 1×10^{-1} Torr

Growth rate:

2.5 nm/sec

The surface of the epitaxial layer was thermally oxidized to a depth of 50 nm. On the thermally oxidation film, a fused silica substrate having been optically polished was superposed and the combined substrates were heated in an oxygen atmosphere at 400 °C for 20 hours, whereby the two substrates were bonded tightly.

The bonded substrates were selectively etched by immersion in a mixture of hydrofluoric acid and aqueous hydrogen peroxide solution (1:5) with stirring. In 62 minutes, the porous Si substrate was selectively etched and removed completely with the monocrystalline Si left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the above etching solution is as low as about 20 $\mbox{\normalfont\AA}$ or a little less for 62 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10^5 or more, and the decrease of the film thickness of the nonporous layer (several tests of $\mbox{\normalfont\AA}$) by etching was practicably negligible.

Consequently, the porous Si substrate of 200 μm thick was removed, and a monocrystalline Si layer of 5 μm thick was formed on the fused silica substrate.

The fused silica substrate having the monocrystalline Si layer was heat-treated at 1000 °C in a hydrogen atmosphere at 760 Torr. The treated sample was evaluated for surface flatness by interatomic microscope observation. The surface roughness before the hydrogen treatment of 20 nm was improved to 1.6 nm.

The resulting Si layer was found to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Example 18

A monocrystallin P-type (l00) Si substrate of 200 μ m thick was anodized in 50% HF solution at a current density of 100 mA/cm². Th porous structur was formed at a rate of 8.4 μ m/min. The entire

15

20

30

45

of th P-type (100) Si substrate of 200 μ m thick was made porous in 24 minut s.

On the porous P-type (100) Si substrate, an pitaxial Si lay r was grown in a thickness of 1 µm by CVD at a low temperature under the deposition conditions below:

Gas: SiH_4 (0.6 L/min), H_2 (100

t/min)

Temperature:

850 · C

Pressure:

40 Torr

Growth rate:

0.3 µm/min

The surface of the epitaxial layer was thermally oxidized to a depth of 50 nm. On the thermally oxidation film, an optically polished glass substrate having a softening point of about 500°C was superposed and the combined substrates were heated in an oxygen atmosphere at 450°C for 0.5 hour, whereby the two substrates were bonded tightly.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 204 minutes, the porous Si substrate was selectively etched and removed completely with the monocrystalline Si left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the above etching solution is as low as about 40 Å or a little less for 204 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decrease of the film thickness of the nonporous layer (several tests of Å) by etching was practicably negligible.

Consequently, the porous Si substrate of 200 μ m thick was removed, and a monocrystalline Si layer of 1 μ m thick was formed on the glass substrate of a low softening point.

This substrate was heat-treated at 900 °C in a hydrogen atmosphere at 10 Torr. The treated sample was evaluated for surface flatness by interatomic microscope observation. The surface roughness before the hydrogen treatment of 20 nm was improved to 1.7 nm.

The resulting Si layer was found to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Example 19

A monocrystallin P-type (100) Si substrate of 525 μ m thick was anodized in 50% HF solution at a curr nt d nsity of 5 mA/cm². The porous structure was form d at a rat of 1 μ m/min. The P-type (100) Si substrate of 525 μ m thick was made porous to a depth of 20 μ m.

The partially porous substrate was heat-treated in an oxyg in atmosphere at 300 °C for one hour.

On the porous face of the P-type (100) Si substrat, an pitaxial Si layer was grown in a thickness of 1.0 μ m at a low temperature by bias sputtering under the deposition conditions below:

RF frequency:

100 MHz

High frequency power:

600 W

Temperature:

300 · C

Ar gas pressure:

 8×10^{-3} Torr

Growth time:

120 minutes

Target DC bias:

-200 V

Substrate DC bias:

+5 V

On the surface of the epitaxial layer, was formed a silicon oxide layer of 500 nm thick by thermal oxidation. On the thermally oxidation film, another silicon substrate was superposed and the combined substrates were heated in a nitrogen atmosphere at 1000 °C for 2 hours, whereby the two substrates were bonded tightly.

The substrate having been made porous was polished on the backside to remove the nonporous portion of the silicon substrate to uncover the porous layer.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (I0:6:50) without stirring. In 20 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the etching solution is as low as about 40 Å or a little less for 204 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the 20 μm thick Si substrate made porous was removed, and a monocrystalline Si layer of 0.75 μm thick was formed on the 500 nm thick silicon oxide layer of the silicon substrate.

The substrate was heat-treated at 900 °C in a hydrogen atmosphere at I0 Torr. The tr at d sampl was valuated for surfac flatness by int ratomic microscop obs rvation. The surfac roughness b fore th hydrog n tr atm nt of 13 nm was found to be improved to 1.4 nm.

The r sulting Si layer was found to have no crystal defect n wly introduced and have satisfac-

15

20

tory crystallinity by observation of cross-s ction by transmission el ctron microscopy.

Exampl 20

A monocrystalline P-type (I00) Si substrate of 600 μ m thick was anodized in 50% HF solution at a current density of 5 mA/cm². The porous structure was formed at a rate of I μ m/min. The P-type (100) Si substrate of 600 μ m thick was made por us to a depth of 20 μ m.

On the porous face of the P-type (I00) Si substrate, an epitaxial Si layer was grown in a thickness of I0 μ m at a low temperature by liquid-phase growth under the growth conditions below:

Solvent:

Sn

Growth temperature:

900 · C

Growth atmosphere:

Ho

Growth time:

20 minutes

On the surface of the epitaxial Si layer, another monocrystalline silicon substrate was superposed which has a silicon oxide layer of 1 µm thick on the surface, and the combined substrates were heated in an oxygen atmosphere at 700 °C for 5 hours, whereby the two substrates were bonded tightly.

The substrate having been made porous was polished on the backside to remove the nonporous portion of the silicon substrate to uncover the porous layer.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 20 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the etching solution is as low as about 10 Å or less for 20 minutes. Therefore, the relative rate of s lective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the Si substrate made porous was removed, and a monocrystalline Si layer of 10 μ m thick was formed on the silicon substrate.

This substrate was h at-tr ated at 950 °C in a hydrogen atmospher at 80 Torr. The treat d sample was evaluated for surface flatness by interatomic microscope observation. The surface roughn ss before the hydrogen treatment of 20 nm was improved to 1.7 nm.

The r sulting Si lay r was found to hav no crystal d fect newly introduced and hav satisfactory crystallinity by observation of cross-section by transmission I ctron microscopy.

Example 2i

On a P-type (100) SI substrate of 200 μ m thick, an epitaxial Si layer was made to grow to a thickness of 0.5 μ m by CVD under the deposition conditions below:

Reactive gas flow rate:

SiH2Cl2, 1000 SCCM

H₂, 230 t/min

Temperature:

1080 ° C

Pressure:

80 Torr

Time:

1 min

This substrate was anodized in a 50 wt% HF solution at a current density of 100 mA/cm². The porous structure formation rate was 8.4 µm/min. The entire of the P-type (100) Si substrate of 200 µm thick was made porous in 24 minutes. In the anodization, the P-type (100) Si substrate only was made porous as described before, and the Si epitaxial layer remained intact.

The surface of the epitaxial layer was thermally oxidized to a depth of 50 nm. On this thermally oxidized layer, a fused silica substrate which had been optically polished was superposed. The superposed two substrates were heated in an oxygen atmosphere at 800 °C for 3 hours, thereby the two substrates being bonded tightly.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 204 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the etching solution is as low as about 40 Å or less for 204 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10^5 or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the 200 μ m-thick porous Si substrate was removed, and a monocrystallin. Si layer of 0.5 μ m thick was formed on the silicon substrate.

This substrat was heat-treated at 950 °C in a hydrogen atmosphere at 80 Torr. The treated sample was evaluated for surface flatness by interatomic microscope observation. The surface rough-

25

30

35

ness before the hydrogen treatment of 20 nm was improv d to 1.7 nm.

The resulting Si layer was found to have no crystal defect n wly introduced and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Example 22

On the surface of a P-type (I00) Si substrate of 200 μ m thick, an N-type Si layer of 1 μ m thick was formed by proton ion implantation with an H⁻ implantation quantity of 5 \times I0¹⁵ (ions/cm²). This substrate was anodized in 50% HF solution at a current density of 100 mA/cm². The porous structure was formed at a rate of 8.4 μ m/min. The entire of the P-type (100) Si substrate of 200 μ m thick was made porous in 24 minutes. In the anodization, the P-type (100) Si substrate only was made porous with the N-type Si layer remained intact.

Th surface of the N-type monocrystal layer was thermally oxidized to a depth of 50 nm. On the thermally oxidation film, a fused silica substrate having been optically polished was superposed and the combined substrates were heated in an oxygen atmosphere at 800 °C for 0.5 hour, whereby the two substrates were bonded tightly.

The bonded two substrates were covered by deposition of Si_3N_4 in a thickness of 0.1 μm by low-pressure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

Then the bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 204 minutes, the porous Si substrate was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

The rate of etching of the nonporous Si layer by the etching solution is as low as about 40 Å or a little less for 204 minutes. Therefore, the relative rate of selective etching of the porous layer to that of nonporous layer is 10⁵ or more, and the decrease of the film thickness of the nonporous layer (several tens of Å) by etching was practicably negligible.

Consequently, the porous Si substrate of 200 μ m was removed, and after removal of the Si₃N₄ lay r, monocrystalline Si layer of 1.0 μ m thick was formed uniformly with no def ct on th silicon oxide surfac of th silicon substrate.

In the cas $% 10^{10}\,M_{\odot}$ where $% 10^{10}\,M_{\odot}$ has replaced by Api zon Wax coating or Electron Wax coating, the effect was the same, and the porous Si substrate only was completely removed.

This Si substrate was h at-treated at 950 °C in a hydrogen atmosphere at 80 Torr. The tr ated sample was evaluated for surface flatness by interatomic microscop obs rvation. Th surface roughness before the hydrogen treatment of 20 nm was improved to 1.7 nm.

The resulting Si layer was found to have no crystal defect newly introduced and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Example 23

A monocrystalline P-type (I00) Si substrate of 200 μm thick was anodized in 50% HF solution at a current density of 5 mA/cm². The porous structure was formed at a rate of 0.9 μm/min. The entire of the P-type (100) Si substrate of 200 μm thick was made porous in 223 minutes.

The resulting porous Si substrate was heattreated in a hydrogen atmosphere under the conditions below to obtain a smooth layer on the surface:

Temperature: 950 ° C
Pressure: 80 Torr
Time: 25 minutes

The smooth layer on the surface was observed by high-resolution scanning microscopy, RHEED, and it was found that a thin monocrystal layer of 20 nm thick of the same orientation as that of the substrate.

On the surface of this thin monocrystal layer, another Si substrate was superposed which has an oxide layer of 5000 Å thick, and the combined substrates were heated in an oxygen atmosphere at 800°C for 0.5 hour, thereby the two substrates being bonded tightly.

The bonded two substrates were covered by deposition of Si_3N_4 in a thickness of 0.1 μm by low pressure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (I0:6:50) without stirring. In 205 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material.

After removal of the Si_3N_4 layer, a thin monocrystalline Si layer was formed on the SiO_2 .

The resulting Si lay r was confirmed to have no crystal d fect n wly introduc d and have satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

50

10

15

30

35

40

45

50

55

Exampl 24

A monocrystalline P-type (I00) Si substrate of 200 μ m thick was anodized in 50% HF solution at a current density of 100 mA/cm². The porous structur was formed at a rate of 8.4 μ m/min. The entire of the P-type (I00) Si substrate of 200 μ m thick was made porous in 24 minutes.

The resulting porous Si substrate was heattreated in a hydrogen atmosphere under the conditions below to obtain a smooth layer on the surface:

Temperature:

950 ° C

Pressure:

50 Torr

Time:

45 minutes

Th smooth layer on the surface was observed by high-resolution scanning microscopy, RHEED, and it was found that a thin nonporous monocrystal layer of 50 nm thick of the same orientation as that of th substrate.

This epitaxial layer was oxidized in a thickness of 10 nm. On the surface of the oxide layer, another Si substrate was superposed which has an oxide layer of 5000 Å thick, and the combined substrates were heated in an oxygen atmosphere at 900 °C for 0.5 hour, thereby the two substrates being bonded tightly.

The bonded two substrates were covered by deposition of Si_3N_4 in a thickness of 0.1 μm by low pressure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were etched by immersion in 6M KOH solution. As described before, the etching rate of the ordinary monocrystalline Si by KOH is about 1 μm or less per minute, while the etching rate of the porous Si layer is about 100 times that of the nonporous monocrystalline Si. Therefore, the porous Si substrate of 200 μm thick was removed in 2 minutes. After removal of the Si₂N₄, a monocrystalline Si was formed with excellent crystallinity on the SiO₂.

Example 25

A monocrystalline P-type (100) Si substrate of 200 μ m thick was anodized in 50% HF solution at a current density of l00 mA/cm². The porous structure was formed at a rate of 8.4 μ m/min. The entire of the P-type (l00) Si substrate of 200 μ m thick was made porous in 24 minutes.

The r sulting porous Si substrat was washed with I.5% hydrofluoric acid solution, and then imm diat ly h at-treated in an argon atmospher und r th conditions below to obtain a smooth lay r on the surface:

T mp rature: 950 °C

Pressure: 1 Torr

Time:

60 minutes

The smooth layer on the surface was observed by high-resolution scanning microscopy, RHEED, and it was found that a thin monocrystal layer of 20 nm thick was formed in the same orientation as the substrate.

On the surface of this thin monocrystal layer, another Si substrate was superposed which has an oxide layer of 5000 Å thick, and the combined substrates were heated in an oxygen atmosphere at 800 °C for 0.5 hour, thereby the two substrates being bonded tightly.

The bonded two substrates were covered by deposition of Si_3N_4 in a thickness of 0.1 μ m by low pressure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were etched by immersion in a hydrofluoric nitric acetic acid solution (I:3:8). As described above, the etching rate of the ordinary monocrystalline Si by the hydrofluoric nitric acetic acid is about 1 μ m or less per minute, while the etching rate of thee porous Si layer is about 100 times that of the nonporous monocrystalline Si. Therefore, the porous Si substrate of 200 μ m thick was removed in 2 minutes. After removal of the Si₃N₄, a monocrystalline Si layer was formed on the SiO₂.

The same effect of selective and complete removal of the porous Si substrate could be obtained when Apiezon Wax was used for the covering in place of the Si₃N₄ layer.

Example 26

A monocrystalline P-type (100) Si substrate of 300 μ m thick was anodized in 50% HF solution at a current density of 5 mA/cm². The porous structure was formed at a rate of 0.9 μ m/min. The surface of the P-type (100) Si substrate of 300 μ m thick was made porous to a thickness of 30 μ m.

The resulting partly porous SI substrate was heat-treated in a hydrogen atmosphere under the conditions below to obtain a smooth layer on the surface:

Temperature:

950 · C

Pressure: Time: 60 Torr 25 minutes

The smooth layer on the surface was observed by high-resolution scanning microscopy, RHEED, and it was found that a thin monocrystal layer of 30 nm thick was formed in the same orientation as that of the substrat

This monocrystalline lay r was oxidized to a thickness of 100 Å. Onto the surface of the oxide lay r, anoth r Si substrate was attached clos ly which has an oxide layer of 500 Å thick, and th combined substrates wer heated in an oxygen

35

40

50

atmospher at 700 °C. for 0.5 hour, th reby th two substrates being bonded tightly.

Th partly porous Si substrat was abraded from the nonporous side to r mov 275 µm ther of by a usual wafer lapping method to uncover the porous silicon.

The substrates were covered by deposition of Si₂N₄ in a thickness of 0.1 μm by low pressure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 30 minutes, the p rous Si substrate only was selectively etched and removed completely with the monocrystalline Si lay r left unetched as an etch-stop material. Aft r r moval of the Si₃N₄ layer, a monocrystalline Si layer was formed on the SiO₂.

Th same effect of selective and complete removal of the porous Si substrate could be obtained when Apiezon Wax was used for the covering in place of the Si₃N₄ layer.

Example 27

A monocrystalline P-type (I00) Si substrate of 200 µm thick was anodized in 50% HF solution at a current density of 100 mA/cm². The porous structure was formed at a rate of 8.4 µm/min. The entire of th P-type (100) Si substrate of 200 µm thick was made porous in 24 minutes.

The resulting porous Si substrate was heattreated in a hydrogen atmosphere under the conditions below to obtain a smooth layer on the surfac :

Temperature:

950 ° C

Pr ssure:

760 Torr

Time:

80 minutes

The smooth layer on the surface was observed by high-resolution scanning microscopy, RHEED, and it was found that a thin monocrystal layer of 20 nm thick was formed in the same orientation as that of th substrate.

On the surface of this thin monocrystal layer, another Si substrate was attached closely which has an oxide layers of 5000 Å thick, and the combined substrates were heated in an oxygen atmosphere at 700 °C for 0.5 hour, thereby the two substrates being bonded tightly.

Th bonded two substrates were covered by deposition of Si₃N₄ in a thickness of 0.1 µm by low pr ssure CVD. Th n th nitride film on th porous substrat only was removed by r active ion etching.

The bonded substrates were etched by imm rsion in a hydrofluoric nitric acetic acid solution (1:3:8). As d scribed above, the etching rate of the

ordinary monocrystalline Si by the hydrofluoric nitric acetic acid is about I µm or less per minute, while the etching rate of the porous Si lay r is about 100 times that of th nonporous monocrystalline Si. Therefore, the porous Si substrate of 200 µm thick was removed in 2 minutes. After removal of the Si₂N₄, a monocrystalline Si layer was formed on the SiO2.

Subsequently, a monocrystalline Si layer was epitaxially grow in a thickness of 2 µm from the thin monocrystal layer by ordinary CVD. The growth conditions are as below.

Gas:

SiH2Ck/H2 1/180 (1/min)

Temperature:

1080 ° C

Pressure: 80 Torr

As the result, a monocrystalline Si layer of 2 um thick was formed on the SiO2.

Example 28

A monocrystalline P-type (I00) Si substrate of 200 µm thick was anodized in 50% HF solution at a current density of 100 mA/cm². The porous structure was formed at a rate of 8.4 µm/min. The entire of the P-type (100) Si substrate of 200 µm thick was made porous in 24 minutes.

The resulting porous Si substrate was washed with 1.5% hydrofluoric acid solution, end then immediately heat-treated in a vacuum chamber under the conditions below to obtain a smooth layer on the surface:

Temperature:

950 °C

Pressure:

 1×10^{-8} Torr

Time:

100 minutes

The smooth layer on the surface was observed by high-resolution scanning microscopy, RHEED, and it was found that a thin monocrystal layer of 15 nm thick was formed in the same orientation as that of the substrate.

On the surface of this thin monocrystal layer, another Si substrate was superposed which has an oxide layer of 5000 Å thick, end the combined substrates were heated in an oxygen atmosphere at 800 °C for 0.5 hour, thereby the two substrates being bonded tightly.

The bonded two substrates were covered by deposition of Si₃N₄ in a thickness of 0.l μm by low pressure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etch-

The bonded substrates were selectively etched by immersion in a mixtur of buff r d hydrofluoric acid, alcohol, and aqu ous hydrogen peroxid solution (10:6:50) without stirring. In 205 minutes, th porous Si substrat only was selectiv ly etch d and removed completely with the monocrystalline Si layer I ft unetched as an etch-stop material. Aft r removal of the Si₃N₄ layer, a thin mon-

25

35

40

45

50

55

ocrystalline Si layer was form d on the SiO2.

The resulting Si layer was confirmed to have no crystal defect newly introduced and keep satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Example 29

A monocrystalline P-type (l00) Si substrate of 200 μ m thick was anodized in 50% HF solution at a current density of 5 mA/cm². The porous structure was formed at a race of 0.9 μ m/min. The entire of the P-type (100) Si substrate of 200 μ m thick was made porous in 223 minutes.

The resulting porous Si substrate was heattreated in a hydrogen atmosphere under the conditions below to obtain a smooth layer on the surface:

Temperature:

950 ° C

Pressure:

80 Torr

Time:

25 minutes

The smooth layer on the surface was observed by high-resolution scanning microscopy, RHEED, and it was found that a thin monocrystal layer of 20 nm thick of the same orientation as that of the substrat.

On the surface of this thin monocrystal layer, an optically polished fused silica substrate was superposed, and the combined substrates were heated in an oxygen atmosphere at 800 °C for 0.5 hour, thereby the two substrates being bonded tightly.

The bonded two substrates were covered by deposition of Si_3N_4 in a thickness of 0.1 μm by low pressure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were selectively etched by immersion in a mixture of 49% hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 65 minutes, the porous SI layer only was selectively etched and r moved completely with the monocrystalline Si lay r left unetched as an etch-stop material. After removal of the Si₂N₄ layer, a thin monocrystalline SI layer was formed on the fused silica substrate.

The resulting Si layer was confirmed to have no crystal defect newly introduced and keep satisfactory crystallinity by observation of cross-section by transmission electron microscopy.

Example 30

A monocrystalline P-type (100) Si substrat of 200 μ m thick was anodized in 50% HF solution at a current density of 100 mA/cm². The porous structure was formed at a rate of 8.4 μ m/min. The entire of th P-type (100) Si substrat of 200 μ m thick

was made porous in 24 minutes.

The resulting porous SI substrate was heattreated in a hydrog n atmosph r und r the conditions below to obtain a smooth lay r on th surface:

Temperature:

950 ° C

Pressure:

50 Torr

Time:

45 minutes

The smooth layer on the surface was observed by high-resolution scanning microscopy, RHEED, and it was found that a thin nonporous monocrystal layer of 50 nm thick of the same orientation as that of the substrate.

This epitaxial layer was oxidized to a thickness of 10 nm. On the surface of the oxide layer, an optically polished glass substrate having the softening point of about 500 °C was superposed, and the combined substrates were heated in an oxygen atmosphere at 450 °C for 0.5 hour, thereby the two substrates being bonded tightly.

The bonded two substrates were covered by deposition of Si_3N_4 in a thickness of 0.1 μ m by low pressure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were etched by immersion in 6M KOH solution. As described before, the etching rate of the ordinary monocrystalline Si by KOH is about 1 μ m or less per minute, while the etching rate of the porous Si layer is about 100 times that of the nonporous monocrystalline Si. Therefore, the porous Si substrate of 200 μ m thick was removed in 2 minutes. After removal of the Si₃N₄, a monocrystalline Si was formed with excellent crystallinity on the low-softening-point glass. substrate.

Example 31

A monocrystalline P-type (I00) Si substrate of 200 μ m thick was anodized in 50% HF solution at a current density of I00 mA/cm². The porous structure was formed at a rate of 8.4 μ m/min. The entire of the P-type (100) Si substrate of 200 μ m thick was made porous in 24 minutes.

The resulting porous Si substrate was washed with I.5% hydrofluoric acid solution, and then immediately heat-treated in an argon atmosphere under the conditions below to obtain a smooth layer on the surface:

Temperature:

950 ° C

Pressur :

60 minut s

Th smooth layer on the surface was obs rv d by high-resolution scanning microscopy, RHEED, and it was found that a thin monocrystal layer of 20 nm thick was formed in the same orientation as that of the substrate.

On the surface of this thin monocrystal layer, an optically polish diglass substrate having the softening point of about 500 °C was superposed, and the combined substrates were high an oxygen atmosphere at 450 °C for 0.5 hour, thereby the two substrates being bonded tightly.

Th bonded two substrates were covered by deposition of Si_2N_4 in a thickness of 0.1 μm by low pr ssure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were etched by immersion in a hydrofluoric nitric acetic acid solution (1:3:8). As described above, the etching rate of the ordinary monocrystalline Si by the hydrofluoric nitric acetic acid is about 1 μ m or less per minute, whil the etching rate of the porous Si layer is about 100 times that of the nonporous monocrystalline Si. Therefore, the porous Si substrate of 200 μ m thick was removed in 2 minutes. After removal of the Si₃N₄ layer, a monocrystalline Si layer was formed on the low-softening point glass substrate.

The same effect of selective and complete removal of the porous Si substrate could be obtained when Apiezon Wax was used for the covering in place of the Si₃N₄ layer.

Example 32

A monocrystalline P-type (I00) Si substrate of 300 μ m thick was anodized in 50% HF solution at a current density of 5 mA/cm². The porous structure was formed at a rate of 0.9 μ m/min. The surfac of the P-type (100) Si substrate of 300 μ m thick was made porous to a thickness of 30 μ m.

The resulting partly porous Si substrate was heat-tr ated in a hydrogen atmosphere under the conditions below to obtain a smooth layer on the surface:

Temperature:

950 °C

Pressure:

60 Torr

Time:

25 minutes

Th smooth layer on the surface was observed by high-r solution scanning microscopy, RHEED, and it was found that a thin monocrystal layer of 30 nm thick was formed in the same orientation as that of the substrate.

This monocrystalline layer was oxidized to a thickness of 100 Å. Onto the surface of the oxide layer, a fused silica substrate was attached closely, and the attached substrat s w r h at d in an oxygen atmosphere at 700 °C for 0.5 hour, th reby the two substrates being bond d tightly.

The partly porous Si substrat was abrad d from the nonporous side to remove 270 µm thereof by a usual wafer lapping method to uncover the porous silicon.

The substrates were cov red by d position of Si_3N_4 in a thickness of 0.1 μm by low pr ssur CVD. Then the nitride film on the porous substrationly was removed by reactive ion teching.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (10:6:50) without stirring. In 30 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material. After removal of the Si₂N₄ layer, a monocrystalline Si layer was formed on the fused silica substrate.

The same effect of selective and complete removal of the porous Si substrate could be obtained when Apiezon Wax was used for the covering in place of the Si₃N₄ layer.

Example 33

20

25

A monocrystalline P-type (100) Si substrate of 200 μ m thick was anodized in 50% HF solution at a current density of 100 mA/cm². The porous structure was formed at a rate of 8.4 μ m/min. The entire of the P-type (100) Si substrate of 200 μ m thick was made porous in 24 minutes.

The resulting porous Si substrate was heattreated in a hydrogen atmosphere under the conditions below to obtain a smooth layer on the surface:

Temperature:

950 · C

Pressure:

760 Torr

Time:

760 Forr 80 minutes

The smooth layer on the surface was observed by high-resolution scanning microscopy, RHEED, and it was found that a thin monocrystal layer of 20 nm thick was formed in the same orientation as that of the substrate.

On the surface of this thin monocrystal layer, an optically polished glass substrate having the softening point of about 800 °C was attached closely, and the attached substrates were heated at 700 °C for 0.5 hour, thereby the two substrates being bonded tightly.

The bonded two substrates were covered by deposition of Si₃ N₄ in a thickness of 0.1 μm by low pressure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were etched by immersion in a hydrofluoric nitric acetic acid solution (1:3:8). As described above, the tching rate of th ordinary monocrystallin. Si by the hydrofluoric nitric ac tic acid is about 1 μ m or 1 ss per minute, while the tching rate of the porous Si layer is about 100 times that of the nonporous monocrystalline Si. Therefore, the perious Si substrate of 200 μ m thick was removed in 2 minutes. After

20

30

35

40

45

50

55

removal of the Si_3N_4 , a monocrystalline Si layer was formed on the glass substrat .

Subsequently, a monocrystalline Si lay r was epitaxially grown in a thickn ss of 2 μm from the thin monocrystal layer by ordinary CVD. The growth conditions are as below.

Gas:

SiH₂Cl₂/H₂ 1/180 (1/min)

T mperature:

1080 °C

Pressure: 80 Torr

As the result, a monocrystalline Si layer of 2 μ m thick was formed on the glass substrate.

Example 34

A monocrystalline P-type (I00) Si substrate of 200 μ m thick was anodized in 50% HF solution at a current density of 100 mA/cm². The porous structur was formed at a rate of 8.4 μ m/min. The entire of the P-type (I00) Si substrate of 200 μ m thick was made porous in 24 minutes.

The resulting porous Si substrate was washed with 1.5% hydrofluoric acid solution, and then immediately heat-treated in a vacuum chamber under the conditions below to obtain a smooth layer on the surface:

Temperature:

950 · C

Pressure: Time: 1×10^{-8} Torr

71. -

100 minutes

The smooth layer on the surface was observed by high-resolution scanning microscopy, RHEED, and it was found that a thin monocrystal layer of 15 nm thick was formed in the same orientation as that of the substrate.

On the surface of this thin monocrystal layer, an optically polished fused silica substrate was superposed, and the combined substrates were heated in an oxygen atmosphere at 800 °C for 0.5 hour, thereby the two substrates being bonded tightly.

The bonded two substrates were covered by deposition of Si_3N_4 in a thickness of 0.1 μm by low pressure CVD. Then the nitride film on the porous substrate only was removed by reactive ion etching.

The bonded substrates were selectively etched by immersion in a mixture of buffered hydrofluoric acid, alcohol, and aqueous hydrogen peroxide solution (I0:6:50) without stirring. In 205 minutes, the porous Si substrate only was selectively etched and removed completely with the monocrystalline Si layer left unetched as an etch-stop material. After removal of th Si₃N₄ lay r, a thin monocrystalline Si layer was form d on the fused silica substrate.

The r sulting Si layer was confirm d to hav no crystal def ct newly introduced and keep satisfactory crystallinity by observation of cross-section by transmission electron microscopy. The same ffect of s lective and complet removal of the porous Si substrat could be obtained when Api zon Wax was used for the covering in place of the Sl_3N_4 layer.

As described in detail above, the present invention provides an excellent process for producing on an insulating substrate a crystalline Si layer having high crystallinity corresponding to monocrystalline wafer, which is superior in productivity, uniformity, controllability, and cost.

The present invention provides a process for producing a semiconductor substrate which has the advantages of conventional SOI devices and is applicable to various high-performance electronic devices, integrated circuits and the like.

The present invention provides a process for producing a semiconductor substrate which can replace the expensive SOS and SIMOX in production of large-scale integrated circuits of SOI structure.

A process for producing a semiconductor substrate comprises heat-treating a substrate having a monocrystal thereon in a reducing atmosphere.

25 Claims

- A process for producing a semiconductor substrate, comprising heat-treating a substrate having a monocrystal thereon in a reducing atmosphere.
- A process for producing a semiconductor substrate according to Claim 1, wherein the reducing atmosphere is an atmosphere containing hydrogen.
- A process for producing a semiconductor substrate according to Claim 1 or Claim 2, wherein the heat treating in the reducing atmosphere is conducted at a pressure not higher than atmospheric pressure.
- 4. A process for producing a semiconductor substrate according to any of Claims 1 to 3, wherein the monocrystal is silicon.
- 5. A process for producing a semlconductor substrate, comprising the steps of porousifying a silicon base member, forming a nonporous monocrystalline silicon layer on the porous material to prepare a first substrate, heat-treating the first substrate at a temperature lower than the melting point of the nonporous monocrystalline silicon in a reducing atmosphere; bonding the surface of the nonporous monocrystalline silicon layer onto a second substrate with interpesition of an insulating layer, and the normoving the porous silicon by etch-

15

25

30

35

45

50

55

ing.

- A process for producing a semiconductor substrate according to Claim 5, wher in the reducing atmosphere is a hydrogen atmosphere.
- 7. A process for producing a semiconductor substrate according to Claim 5, wherein the heat treating in the reducing atmosphere is conducted at a pressure not higher than atmospheric pressure.
- 8. A process for producing a semiconductor substrate according to Claims 5, wherein the first substrate is formed through the step of porousifying a silicon base member, and the step of forming a monocrystalline silicon layer on the porous silicon.
- A process for producing a semiconductor substrate according to Claims 5, wherein the insulating layer is a silicon oxide layer.
- 10. A process for producing a semiconductor substrate according to Claims 5, wherein the porous silicon is selectively removed by etching with an HF-containing solution.
- 11. A process for producing a semiconductor substrate according to Claims 5, wherein the monocrystalline silicon layer formed on the porous silicon substrate has a thickness of not more than 20 µm.
- 12. A process for producing a semiconductor substrate according to Claims 5, wherein the second substrate is a silicon substrate.
- 13. A process for producing a semiconductor substrate according to Claims 5, wherein the second substrate is a light-transmissive substrate.
- 14. A process for producing a semiconductor substrate according to Claims 5, wherein the step of the bonding includes operation in an oxygen-containing atmosphere.
- 15. A process for producing a semiconductor substrate according to Claims 5, wherein the step of the bonding includes operation in a nitrogen-containing atmosphere.
- 16. A proc ss for producing a semiconductor substrat, comprising the steps of porousifying a silicon base member, forming a nonporous monocrystalline silicon layer on the porous material to prepar a first substrate, bonding the surface of the nonporous monocrystalline.

silicon onto a second substrate with interposition of an insulating layer, removing the porous silicon by tching, and heat-tr ating the bonded substrates at a temp rature lower than the melting point of the nonporous monocrystalline silicon in a reducing atmosphere.

- 17. A process for producing a semiconductor substrate according to Claim 16, wherein the reducing atmosphere is a hydrogen atmosphere.
- 18. A process for producing a semiconductor substrate according to Claim 16, wherein the heat treating in the reducing atmosphere is conducted at a pressure not higher than atmospheric pressure.
- 19. A process for producing a semiconductor substrate according to Claims 16, wherein the first substrate is formed through the step of porousifying a silicon base member, and the step of forming a monocrystalline silicon layer on the porous material.
- 20. A process for producing a semiconductor substrate according to Claims 16, wherein the insulating layer is a silicon oxide layer.
- 21. A process for producing a semiconductor substrate according to Claims 16, wherein the porous silicon is selectively etched with an HFcontaining solution.
- 22. A process for producing a semiconductor substrate according to Claims 16, wherein the monocrystalline silicon layer formed on the porous silicon substrate has a thickness of not more than 20 μm.
- 23. A process for producing a semiconductor substrate according to Claims 16, wherein the second substrate is a silicon substrate.
 - 24. A process for producing a semiconductor substrate according to Claims 16, wherein the second substrate is a light-transmissive substrate.
- 25. A process for producing a semiconductor substrate according to Claims 16, wherein the step of the bonding includes operation in an oxyg n-containing atmosph r.
- 26. A proc ss for producing a s miconductor substrate according to Claims 16, wherein the step of the bonding includes op ration in a nitrog n-containing atmosph re.

27. A process for producing a semiconductor substrate according to Claims 16, wherein th nonporous monocrystalline silicon lay r is formed through epitaxial growth.

28. A process for producing a semiconductor substrate, comprising the steps of porousifying a silicon substrate, heat-treating the porousified silicon substrate at a temperature lower than the melting point thereof in a reducing atmosphere to change the surface layer of the porousified silicon substrate into a nonporous monocrystalline silicon layer; bonding the nonporous monocrystalline silicon layer onto the surface comprising an insulating material of a and removina second substrate. porousified portion of the silicon substrate by etching.

29. A process for producing a semiconductor substrate according to Claim 28, further comprising, subsequent to the etching step, a step of growing a monocrystalline silicon layer though epitaxial growth from the nonporous monocrystalline silicon layer.

30. A process for producing a semiconductor substrate according to Claim 28, wherein the silicon substrate to be made porous is of a Ptype.

31. A process for producing a semiconductor substrate according to Claims 28, wherein the step of the bonding includes operation in an oxygen-containing atmosphere.

32. A process for producing a semiconductor substrate according to Claims 28, wherein the step of the bonding includes operation in a nitrogen-containing atmosphere.

33. A process for producing a semiconductor substrate according to Claim 28, wherein the reducing atmosphere is an atmosphere containing hydrogen. 5

10

15

20

25

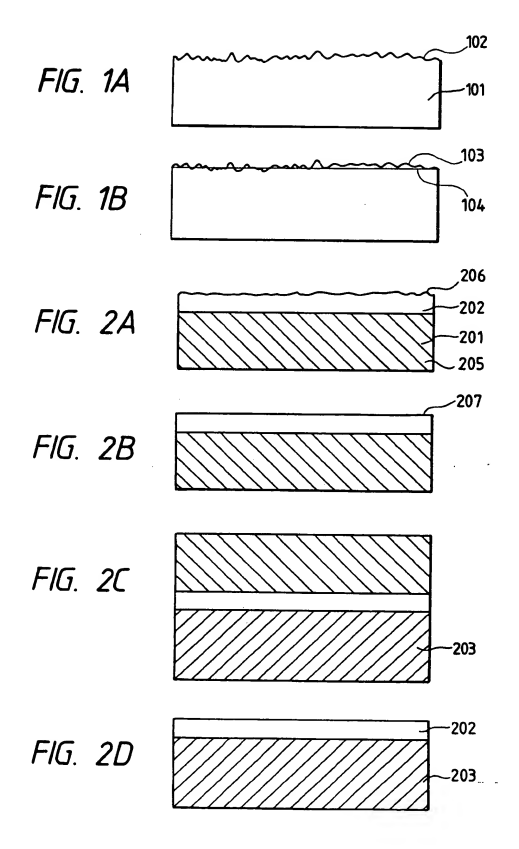
30

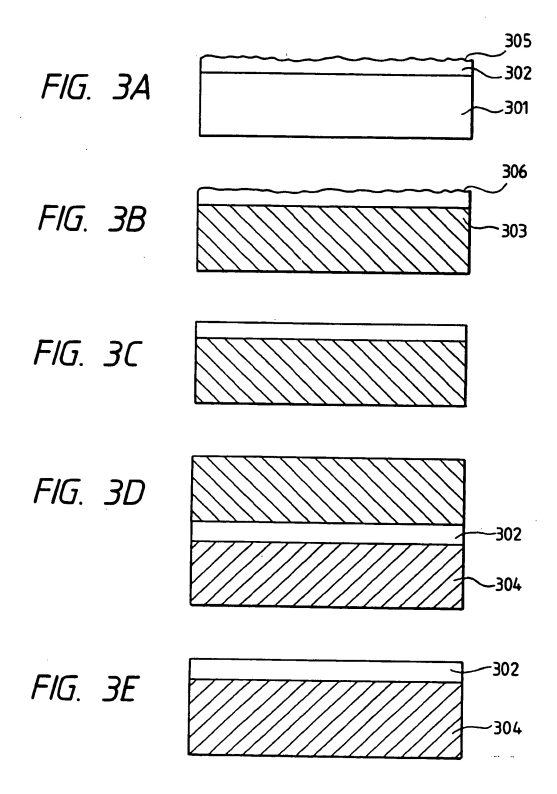
35

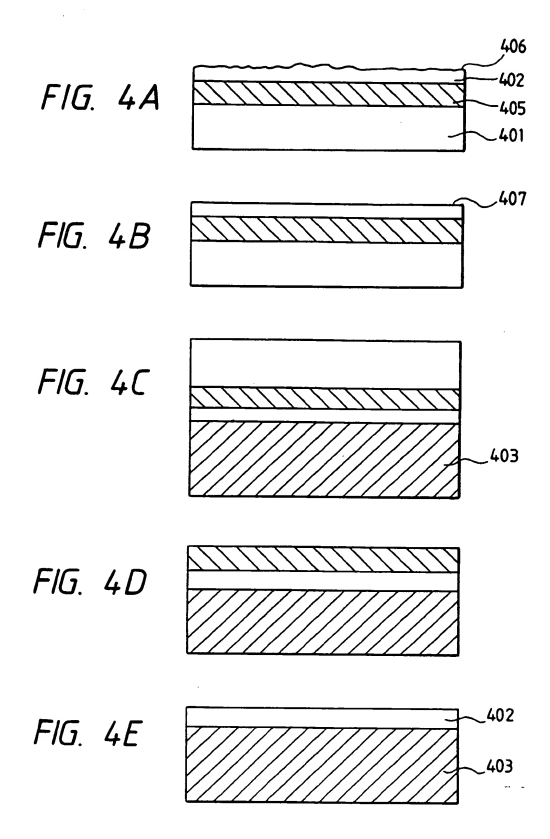
40

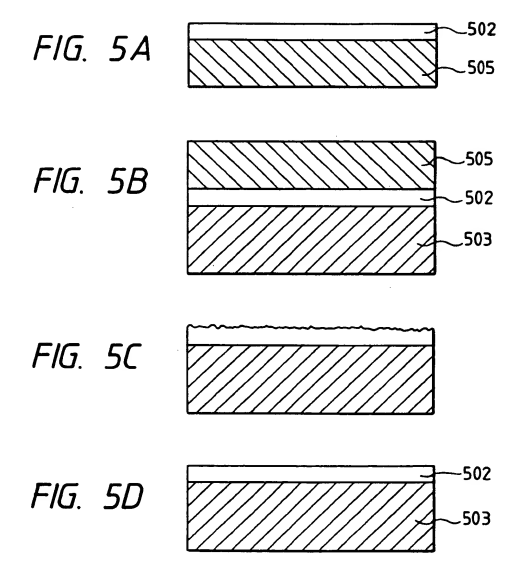
45

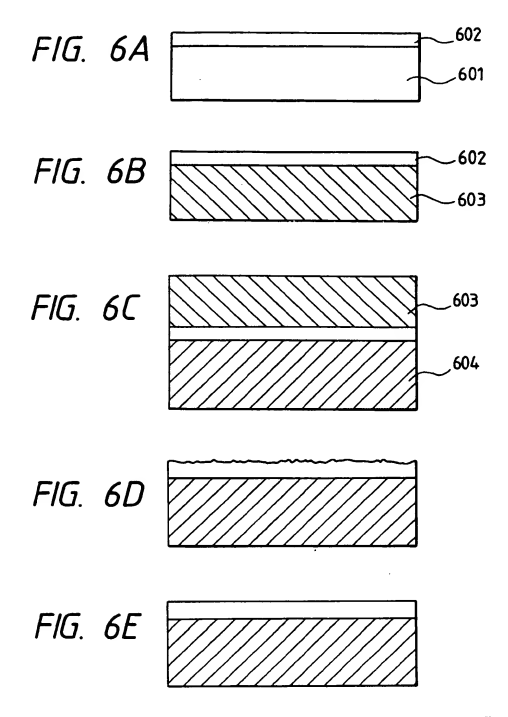
50

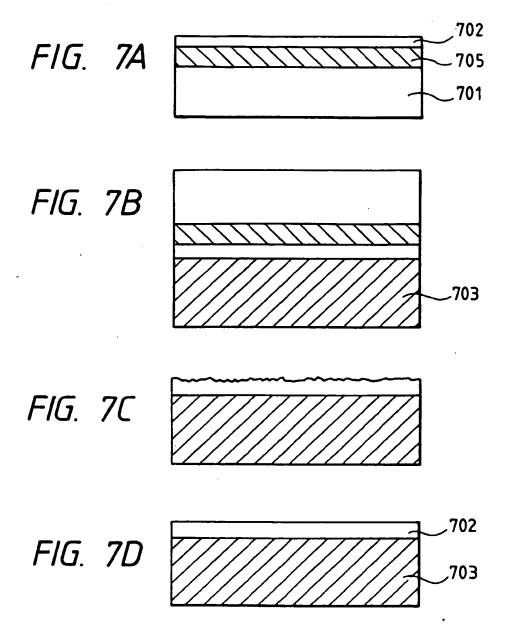


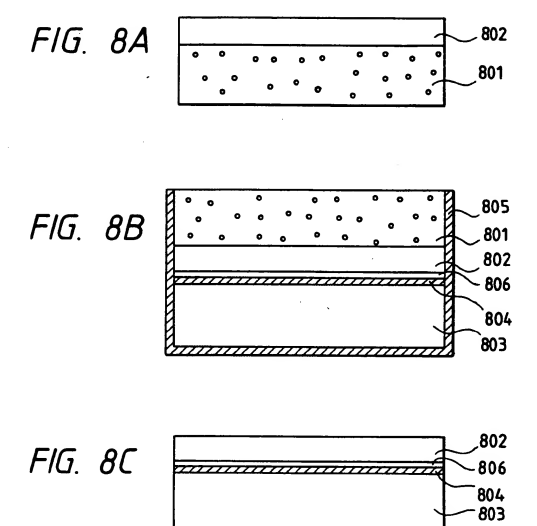


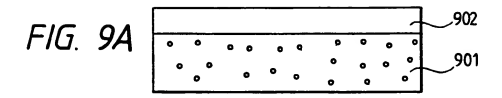


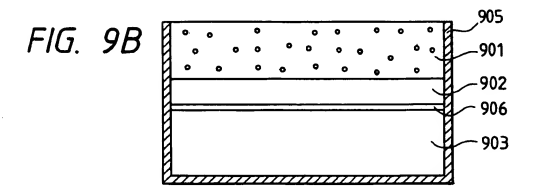












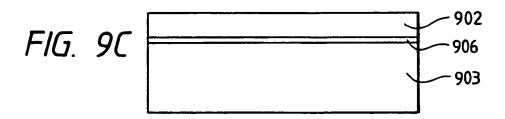


FIG. 10

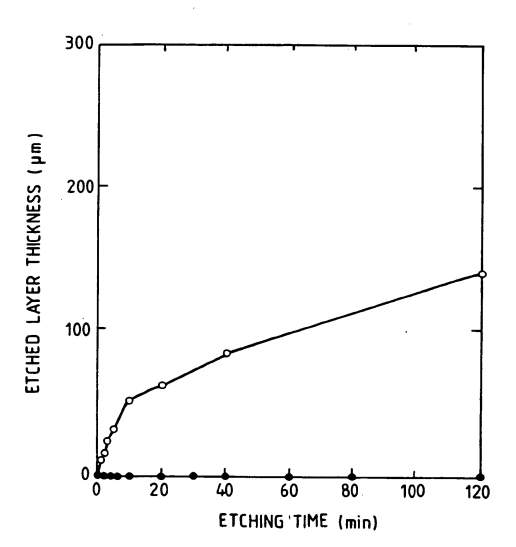
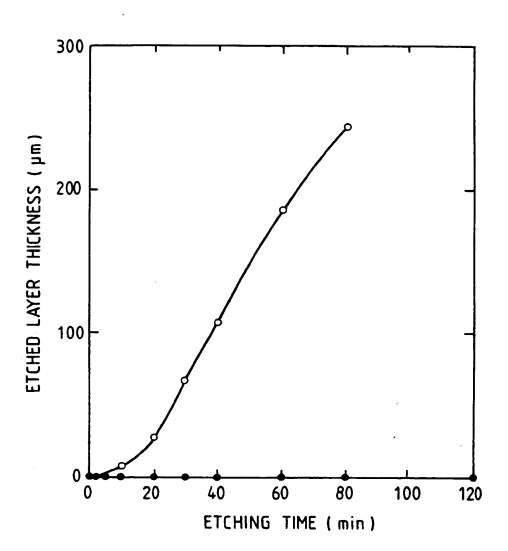
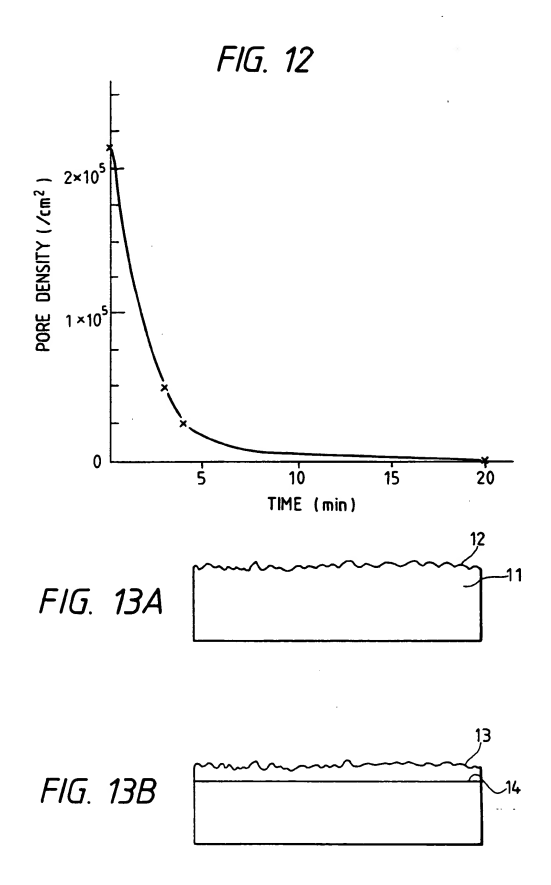


FIG. 11







Europäisches Patentamt

European Patent Office

Office européen des brevets



(11) EP 0 553 852 A3

(12)

EUROPEAN PATENT APPLICATION

(88) Date of publication A3: 14.01.1998 Bulletin 1998/03

(51) Int. Ci.⁶: **H01L 21/76**, H01L 21/20, H01L 21/324

(43) Date of publication A2: 04.08.1993 Bulletin 1993/31

(21) Application number: 93101413.8

(22) Date of filing: 29.01.1993

(84) Designated Contracting States: DE FR GB

(30) Priority: 30.01.1992 JP 38458/92 30.01.1992 JP 38460/92 31.01.1992 JP 16513/92 31.01.1992 JP 16523/92 31.01.1992 JP 16524/92

(71) Applicant: CANON KABUSHIKI KAISHA Tokyo (JP)

(72) Inventors:
Sato, Nobuhlko,
c/o Canon Kabushiki Kaisha
Tokyo 146 (JP)

 Yonehara, Takao, c/o Canon Kabushiki Kaisha Tokyo 146 (JP)

 Sakaguchi, Kiyofumi, c/o Canon Kabushiki Kaisha Tokyo 146 (JP)

(74) Representative:
Tledtke, Harro, Dipl.-Ing.
Patentanwaltsbüro
Tledtke-Bühling-Kinne & Partner
Bavariaring 4
80336 München (DE)

(54) Process for producing semiconductor substrate

(57) A process for producing a semiconductor substrate comprises heat-treating a substrate having a monocrystal thereon in a reducing atmosphere.



EUROPEAN SEARCH REPORT

Application Number EP 93 10 1413

Category	Citation of document with indication, where appropriate, of relevant passages		Relevant	CLASSIFICATION OF THE APPLICATION (IntCL5)
x	OHKURA A ET AL: "The optimization of in-situ thermal cleaning focused on surface microroughness for future Si epitaxial growth", EXTENDED ABSTRACTS OF THE 1991 INTERNATIONAL CONFERENCE ON SOLID STATE DEVICES AND MATERIALS, YOKOHAMA, JAPAN, 27-29 AUG. 1991, 1991, TOKYO, JAPAN, BUS. CENTER ACAD. SOC. JAPAN, JAPAN, PAGE(S) 559 - 561 XP002035569			H01L21/76 H01L21/20 H01L21/324
A	* the whole documer		5,16	
X	faults in a silicor (100) orientation b JOURNAL OF APPLIED	PHYSICS, 1 JUNE 1990, 11, PAGE(S) 7176 - 7178	1,2,4	
A	* the whole documer		5,16	
X	wafer bonding-wafer evaluations", JAPA		1,2,4	TECHNICAL FIELDS SEARCHED (InLCL.5)
A	XP002035570 * page 1436, left-h - paragraph 9 *	and column, paragraph 6	5,16	
P,A	EP θ 515 181 A (CAN * claims 1,17-19 *	10N KK) 25 November 1992	5,27	
A	silicon layer and h JOURNAL OF THE ELEC AUG. 1978, USA, VOL	reat-treatment effect*, TROCHEMICAL SOCIETY, 125, NR. 8, PAGE(S) 0013-4651 XP000616529	28,30,33	
		-/		
	The present search report has			
	Place of search THE HAGUE	Date of completion of the search 11 November 1997	Van	Examiner Craeynest, F
X : parti Y : parti docu A : tech O : non-	ATEGORY OF CITED DOCUMENTS cularly relevant if taken alone cularly relevant if combined with anot ment of the same category nological background written disolosure mediate document	T: theory or principle E: earlier patent door after the filing date her D: dooument cited for L: document ofted for å: rnember of the sau dooument	underlying the in iment, but publis the application r other reasons	ivertion hed on, or



EUROPEAN SEARCH REPORT

Application Numbe

EP 93 10 1413

		ERED TO BE RELEVANT		
ategory	Citation of document with it of relevant pass	ndication, where appropriate, ages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.5)
	on porous silicon"		28,30,33	
	·			TECHNICAL FIELDS SEARCHED (IMLCI.5)
l.	The present search report has b	een drawn up for all claims	-	
	Place of search	Date of completion of the search		Examiner
X : partic Y : partic docur A : techn O : non-	THE HAGUE TEGORY OF CITED DOCUMENTS cularly relevant if taken alone cularly relevant if ormbined with anoth ment of the same category cological background written disclosure mediate document	L : document cited f	le underlying the invoument, but publish te in the application or other reasons	

EPO FORM 1503 03.62 (PO4C01)



Application Number

EP 93 10 1413

CLAIMS INCURRING FEES				
The present European patent application comprised at the time of filing more than ten claims.				
Only part of the claims have been paid within the prescribed time limit. The present European search report has been drawn up for the first ten claims and for those claims for which claims fees have been paid, namely claim(s):				
No claims fees have been paid within the prescribed time limit. The present European search report has been drawn up for the first ten claims.				
LACK OF UNITY OF INVENTION				
The Search Division considers that the present European patent application does not comply with the requirements of unity of invention and relates to several inventions or groups of inventions, namely:				
SEE SHEET B				
All further search fees have been paid within the fixed time limit. The present European search report has been drawn up for all claims. Only part of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the inventions in respect of which search fees have been paid, namely claims: None of the further search fees have been paid within the fixed time limit. The present European search report has been drawn up for those parts of the European patent application which relate to the invention first mentioned in the claims, namely claims:				

- - B-



European Patent Office

LACK OF UNITY OF INVENTION

The Search Division considers that the present European patent application does not comply with the requirement of unity of invention and relates to several inventions or groups of inventions, namely:

- 1. Claims 1-27: Heat treatment of monocrystalline layer on substrate in reducing atmosphere to smoothen its surface
- 2. Claims 28-33: Heat treatment of porous layer on substrate to render it non-porous.